poe60-cys poe60-cys Synthetic Human mat

E.coli op Human KGF

Abp43457 Abb884861 Abb884885 Abb88487 Abb13740 Abb13736 Abb14289 Abb14289 Abb18740 Abb18740 Abb18740 Abb18740 Abb18740 Abb1883 Abb18846 Abb18846 Abb18846 Abb18846 Abb18846 Abb18846 Abb18846 Abb18846 Abb18846

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L. cuprin L. cuprin High mole

Protein e Polythoa

us-09-813-820-4.rag

OM protein -

Run on:

Sequence:

Minimum DB E Maximum DB E

Database

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Result

Searched:

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Collagen binding protein; cna gene; sepsis; infection; microbial surface component regonising adhesive matrix molecule; MSCRAMM; adhesin; vaccine; immunisation; diagnosis; therapy; epitope M31.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Antibody that interacts with collagen binding domain of Staphylococcal cna gene product - useful to prevent bacterial sepsis in animal infected with Staphylococcus aureus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         This protein comprises Staphylococcus aureus collagen binding protein (CBP) epitope M31, i.e. amino acids 61-343 of full-length CBP, plus a
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1. .12
/note= "vector pQE30-derived peptide"
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                                                                                                                                                                                                                                                                                                                           ALIGNMENTS
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                                                       ABUÜB487
ABU13740
ABU13730
ABU13731
ABPS8754
AAB10309
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AAB10709
AAB11871
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/note= "epitope M31"
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N-PSDB; AAT93437.
27-AUG-2003
25-MAR-2003
21-MAY-1998
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Ada 97172 E
Ada 97172 E
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Abu 7901 G
Abu 7901 C
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Compugen Ltd.
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Copyright (c) 1993 - 2004
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Maximum Match 100%
Listing first 45 summaries
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ABM79017
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Symersky J;

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               acid sequences (see AAT93436-38) respectively encode CBP epitopes M17, M31 and M55 (see AAW31552-54) that confer protection against S. aureus infection. These nucleic acid sequences can be used in the recombinant production of the CBP epitopes. The CBP protein and antigenic epitopes are contemplated for use in the treatment of pathological infections, especially to prevent bacterial adheaion to collagen. The claimed nucleic acids as well as claimed anti-CBP antibodies will also be of use in screening, diagnostic and therapeutic applications including active and passive immunisation and methods for the prevention of bacterial
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKI 120
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                                                                                                                                                                            colonisation in an animal such as a human. The CBP epitopes are also contemplated for use in the preparation of vaccines and as carrier proteins in vaccine formulations, as well as in the formulation of compositions for the prevention of S. aureus infection. (Updated on AMAR-2003 to correct PI field.) (Updated on 27-AUG-2003 to correct OS
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/note= "vector pQE30-derived peptide"
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/note= "epitope M17"
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Matches 211, Conservative
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This protein comprises Staphylococcus aureus collagen binding protein (CBP) epitope M55, i.e. amino acids 30-531 of full-length CBP, plus a vector-derived N-terminal peptide. Claimed 441, 849 and 1500 bp nucleic acid sequences (see AAT93436-38) respectively encode CBP epitopes M17, CM 31 and M55 (see AAT93436-38) respectively encode CBP epitopes M17, cm of the CBP epitopes M17, cm of the CBP epitopes The CBP protein and antigenic epitopes are contemplated for use in the treatment of pathological infections, especially to prevent bacterial adhesion to collagen. The claimed nucleic acids as well as claimed anti-CBP antibodies will also be of use in sereening, diagnostic and therapeutic applications including active and consistion in an animal such as a human. The CBP epitopes are also contemplated for use in the preparation of vaccines and as carrier proteins in vaccine formulations, as human. The CBP epitopes are also contemplated for use in the preparation of vaccines and as carrier proteins in vaccine formulations, as well as in the formulation of compensations for the prevention of S. aureus infection. (Updated on 25-KM2-2003 to correct PI field.) (Updated on 27-MUG-2003 to correct OS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 241 KNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVH 300
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                                                                                                                                 Antibody that interacts with collagen binding domain of Staphylococcal cna gene product - useful to prevent bacterial sepsis in animal infected with Staphylococcus aureus.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ----SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 1054.5; DB 2; Length 512; Pred. No. 1.3e-79; 0; Mismatches 0; Indels 115;
                                      <u>ن</u>
                                      Symersky
                                      Sthanam N,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         301 NINANAGIEGTVKGELKVLKQDKDTK 326
                                                                                                                                                                                                                   Claim 31; Page 117-119; 143pp; English
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                                      Hoeoek M, Patti JM, House-Pompeo
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             S. aureus collagen adhesin protein.
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Best Local Similarity 64.7%;
Matches 211; Conservative
(UABR-) UAB RES FOUND.
                                                                             WPI; 1998-008801/01.
                                                                                                  N-PSDB; AAT93438
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Sequence 512 AA;
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Superantigen; SAg; staphylococcal enterotoxin; tumour; cancer; apoptosis; gene therapy; mammalian cell receptor; cytostatic; tumour associated lipid; anergy; T cell; antigen presenting cell; APC; tumouricidal immunocyte; antitumour.

Staphylococcus aureus.

US2002177551-A1.

28-NOV-2002.

30-MAY-2001; 2001US-00870759

31-MAY-2000; 2000US-0208128P

(TERM/) TERMAN D S.

Terman DS;

WPI; 2003-361759/34. N-PSDB; ACA64715. A mammalian cell receptor, useful in the treatment of cancer by binding to tumor associated lipids where the binding induces anergy or apoptosis in T cells and antigen presenting cells.

Disclosure; Page; 167pp; English.

The invention relates to a mammalian cell receptor, useful in the treatment of cancer, which binds to tunnur associated lipids and induces a treatment of cancer, which binds tunnur associated lipids and induces a law included are a mammalian cell useful in the treatment of cancer where the receptor which binds tunnur sasociated lipids and induces to the induces are a tunnucation or death is deleted or functionally deattwated, bridge to concact immunocytes in which they allowing tunnur augmorphism is the probability of a unmonsuppressive fact, and a canadia or salely at unmonsuppressive fact, and a canadia or salely attention of a unmonsuppressive fact, and a construct useful in the treatment of cancer (where an adaptor protein which useful in the treatment of cancer (where an adaptor protein which useful in the treatment of cancer (where an adaptor protein which useful in the treatment of cancer (where an adaptor protein which useful in the treatment of cancer (where an adaptor protein which useful in the treatment of cancer (where an adaptor protein which useful in the treatment of cancer (where some adaptor protein which associated lipids or concact (comprising a lipid raft conjugated to a superantigan) producing (M1) a tunnoricidal immunocyte population useful in the treatment of cancer (comprising a lipid raft conjugated to a superantigan) producing (M2) a tunnoricidal immunocyte population will adaptor protein which adaptor proteins which adaptor at tunnoricidal at mammal (by allowing a tunnoricidal in which adaptor proteins which inhibit T cells on which adaptor proteins which inhibit T cells of administering the tunnoricidally activated T cells on an amamal by allowing a tunnoricidal antigen at administering the tunnoricidally activated T cells on the mammal by allowing a tunnoricidal T cell population in vivo in a mammal (by allowing a tunnoricidal T cell population at vivo in a mammal (by allowing a tunnoricidal T cell population are vivo in a mammal by administering cunnoricidally activated T cells to

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                                                                                                                                                                                                                                     264 TIPQGYGSYNSFSINYKTKITNEQOKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANA 323
or incorporated into a fusion construct with a superantigen. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format from the US patnet office website at "segdata.uspto.gov/sequence.html?DocID=20020177551"
                                                                                                                             12 SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSK
                                                                                                                                                   144 SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSK
                                                                                                                                                                           DITIKDQIQGCQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDV
                                                                                                                                                                                        204 DITIKDQIQGGQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDV
                                                                                                                                                                                                                         TIPOGYGSYNSFSINYKTKITNEQOKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANA
                                                                                                      Gaps
                                                                                                      ;
0
                                                                                Length 1183;
                                                                                                      0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                             CBP; collagen binding protein; mastitis; arthritis.
                                                                               Score 1048; DB 6;
Pred. No. 1.4e-78;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       'note= "claim 11; page 29-30"
                                                                         93.4%; Sco...
100.0%; Pred. No. 1...
0; Mismatches
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/label=_gig_peptide
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                                                                                                                                                                                                                                                                                            GIEGTVKGELKVLKQDKDTK 343
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[180. .1185
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35. .721
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'note= "see CC"
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/label= A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      label= B3
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(first entry)
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                                                                                                                                                                                                                                                                                                                                                                                                                                     Collagen binding protein.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Staphylococcus aureus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               160.
                                                                            Query Match
Best Local Similarity
Matches 200; Conserv
                                                          Sequence 1183 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WO9207002-A1
                                                                                                                                                                                                                                                                                                                                                                                                   25-MAR-2003
04-DEC-1992
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                                                                                                                                                                                                                                                                                                                                                                             AAR22675;
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Integration of the squence serious by the squence assembling a signal sequence. Following this region, a region called A is found a signal sequence. Following this region, a region called A is found a cidal sequence. Following these regions there is a region called BI, B2 and B3. Directly following these regions there is a region called BI, B2 and B3. Consists of a repetitive, hydrophilic structure contg. several proline centides. This region is thought to mediate the binding of the protein to the cell wall. The amino acid sequence nearest to the C-terminal end consists of a long stretch of hydrophobic residues followed by some charged amino acids This region is called M. The CBP can be used for immunisation pref. in combination with a fusion protein, e.g. for vaccination of ruminants against mastitis caused by staphylococcal incertions. It can also be used to block infection in an open skin wound, e.g. for blocking protein receptors or by immunisation. In the latter, the host produces specific antibodies which block the adherence of the bacterial strains to damaged tissue. This treatment can be used for septic arthritis and tissue damage of e.g. skin, connective tissue, and mucous membranes. Dosage for immunisation is 0.5-5 microg CBB/kg; for topical admin. the protein is used at a conc. of 25-250 microg/M1.

(Updated on 25-MAR-2003 to correct PM field.) (Updated on 25-MAR-2003 to
                                                                                                                                                                                                                                                                                                                                                                                                                        (encoded by the sequence assembled from partially
                                                                                                                                                                                                                                                                                                       Hybrid DNA molecule encoding S.aureus collagen binding protein - protein is expressed in B.coli and used for diagnosis e.g. of septic arthritis.
                                                                                                                                                                  Signaes LC;
                                                                                                                                                                  Patti J,
                                                                                                                                                                  Jonsson H, Lindberg KM,
                                                                                                                                                                                                                                                                                                                                                                             Disclosure, Fig 2; 40pp; English.
                                                                                                                 (ALFA ) ALFA LAVAL AGRIC INT AB.
91WO-SE000707
                                             90SE-00003374.
                                                                                                                                                                                                                                   WPI; 1992-167099/20.
N-PSDB; AAQ24123.
                                                                                                                                                                  Hook M,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Sequence 1185 AA;
                                             22-OCT-1990;
22-OCT-1991;
  22-OCT-1991;
                                                                                                                                                                                       Switalski
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17-MAR-2000;
25-APR-2000;
15-AUG-2000;
                                                                                                                                                Hook M,
Domanski
                                                                                                (TEXA )
(UYPA-)
                                                                                    KDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTID 130
                                                                                                                                                                                              VTI PQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINAN 190
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                                                                                                                                                   KDITIKDEIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTID 264
                                                                  DDKVATITSGNKSTNVTVHKSEAGTSSVFYY-KTGDM-LPEDTTHVRWFLNINNEKSYVS 70
                                   Gaps
                                   7
Score 904; DB 2; Length 1185;
Pred. No. 1.5e-66;
3; Mismatches 16; Indels
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                                                                                                                                                                                                                                                                                     AGIEGTVKGELKVLKQDKDTK 345
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 89.68;
                                   Matches 180; Conservative
                  Local Similarity
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Staphylococcus aureus CNA19 protein.

(first entry)

XEXEXEX

RESULT 5 AAE11855

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The invention relates to an antibody which recognises a collagen-binding region including CNA19 of CNA protein from Staphylococcus aureus. This antibody is cross-reactive to collagen binding region of both S. aureus and S. epidermidis. It is useful for preventing or treating S. aureus or S. epidermidis infection in human or animal, and for displacing S. aureus or S. epidermidis bound to collagen. Antibody of the invention is useful for interfering with, modulating, and inhibiting the binding interactions between Staphylococcal bacteria and collagen, for detecting the presence of Staphylococcal bacteria or Staphylococcal collagen or binding proteins, to diagnose Staphylococcal infection, as research tools, for
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      such as
                                                                                                                                                                                                                                                                     .....= "meta strand e, a portion of this region forms a part of the trench in the beta sheet"
89. .92
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                                                                                                                                                                        /note= "Beta strand b; this region forms a part of the trench in the beta sheet"
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                                                                                                                                /note= "Beta strand a; this region forms a part of the
                                                                                                                                                                                                                    /note= "Beta strand d; a portion of this region forms part of the trench in the beta sheet"
82. .84
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  proteins, to diagnose Staphylococcal infection, as research tools, for development of vaccine for passive immunisation against Staphylococcal
                                                                                                                                                                                                                                                                                                                                                                                                                                              /note= "Beta strand h; a portion of this region forms part of the trench in the beta sheet"
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Novel isolated antibody which recognizes collagen-binding peptide eCNA19 peptide from Staphylococcus aureus, useful for preventing or treating Staphylococcus aureus or epidermidis infection.
collagen-binding region; CNA19; Staphylococcus aureus infection; Staphylococcus epidermidis infection; vaccine; CNA protein; cross-reactive antibody.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Casolini F, Patti J,
                                                                                                                                             trench in the beta sheet"
                                                                                                                                                                                                                                                                                                                                "Alpha helix 1"
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110. .115
                                                                                                                                                                                                                                                                                                                                                           "Alpha helix 2"
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/note= "Beta strand j"
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/note= "Beta strand
                                                                                                    Location/Qualifiers
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2000US-0225402P.
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                                                                                                                                                                                                                                                                                                                                                              /note=
                                                                         Staphylococcus aureus
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Ρ;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          infections, and in production facilities or laboratories to isolate additional quantities of collagen-binding proteins. It is also useful for treating medical instruments in order to reduce or eliminate the possibility of them becoming infected or further spreading the infection. The present sequence is S. aureus CNA19 protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Collagen binding protein; ona gene; sepsis; infection; microbial surface component regonising adhesive matrix molecule; MSCRAMM; adhesin; vaccine; immunisation; diagnosis; therapy; epitope M17.
                                                                                                                                                                                                                                                                                                                                                       78
                                                                                                                                                                                                                                                                                                                                                                                                                 9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        This protein comprises Staphylococcus aureus collagen binding protein (CBP) epitope M17, i.e. amino acids 151-297 of full-length CBP, plus a
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           IQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQGYG
                                                                                                                                                                                                                                                                                                                                                       19 ITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIKDQ
                                                                                                                                                                                                                                                                                                                                                                                                              1 ITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIKDQ
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHN 186
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             SYNSFSINYKTKIINEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHN 168
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Symersky
                                                                                                                                                                                                                                       Length 168;
                                                                                                                                                                                                                                                                                              1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /note= "vector pQE30-derived peptide"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Sthanam N,
                                                                                                                                                                                                                                    Score 884; DB 4;
Pred. No. 5.3e-66;
0; Mismatches 1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    House-Pompeo K,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Collagen binding protein M17 epitope.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /note= "epitope M17'
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Claim 31; Page 114; 143pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAW31552 standard; protein; 159 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                UNIV TEXAS A & M SYSTEM
UAB RES FOUND.
                                                                                                                                                                                                                                 78.8%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (revised)
(first entry)
                                                                                                                                                                                                                                                                                           Matches 167; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   13. .159
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (revised)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Staphylococcus aureus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Hoeoek M, Patti JM,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WPI; 1998-008801/01.
                                                                                                                                                                                                                                                                  Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    N-PSDB; AAT93436
                                                                                                                                                                             Sequence 168 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WO9743314-A2
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        14-MAY-1997;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  16-MAY-1996;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              25-MAR-2003
21-MAY-1998
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               20-NOV-1997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        27-AUG-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAW31552;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         139
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        79
                                                                                                                                                                                                                                    Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (UABR-)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Peptide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       with
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caid sequences (see AAT93436-38) respectively encode CBP epitopes M17, and M55 (see AAM1552-54) that confer protection against 8. aureus infection. These nucleic acid sequences can be used in the recombinant production of the CBP epitopes. The CBP protein and antigenic epitopes are contemplated for use in the treatment of pathological infections, capecially to prevent bacterial adhesion to collagen. The claimed nucleic acids as well as claimed anti-CBP antibodies will also be of use in screening, diagnostic and therapeutic applications including active and screening, diagnostic and therapeutic applications including active and connemplated for use in the prevention of bacterial colonisation in an animal such as a human. The CBP epitopes are also contemplated for use in the preparation of vaccines and as carrier proteins in vaccine formulations, as well as in the formulation of S. aureus infection. (Updated on 25-MC-2003 to correct PI field.) (Updated on 27-AUG-2003 to correct OS field.)
                                                                                                                                                                                                                                                                                                                                                                                            1;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      New isolated nucleic acid derived from Enterococcus faecium encoding an Enterococcus faecium polypeptide useful for detection, prevention and treatment of a pathological condition resulting from a bacterial
                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 MRGSHHHHHHGS-----ITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKI
                                                                                                                                                                                                                                                                                                                                                                                                                                1 MRGSHHHHHHGSDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKI
                                                                                                                                                                                                                                                                                                                                                                                              Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Vaccine, urinary tract infection, bacteraemia, endocarditis, wound, abdominal-pelvic infection.
                                                                                                                                                                                                                                                                                                                                                                                              9
                                                                                                                                                                                                                                                                                                                                                      Length 159;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                115 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQA 159
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQA 165
                                                                                                                                                                                                                                                                                                                                                      Score 830; DB 2; Length 15
Pred. No. 1.6e-61;
0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Example 1; SEQ ID NO 6999; 243pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               E. faecium protein sequence SEQ ID 6999.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADC97372 standard; protein; 731 AA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       98US-0085598P.
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                                                                                                                                                                                                                                                                                                                                                          74.0%;
                                                                                                                                                                                                                                                                                                                                                                         96.48;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (first entry)
                                                                                                                                                                                                                                                                                                                                                                                            159; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Enterococcus faecium
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WPI; 2003-799836/75.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Doucette-Stamm LA,
                                                                                                                                                                                                                                                                                                                                                                         Similarity
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                                                                                                                                                                                                                                                                                                                      Sequence 159 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     30-JUN-1998;
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Best Local S
Matches 159
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADC97372;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     RESULT 7
ADC97372
           임
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Entercoccus faccium encoding an Entercoccus faccium polypeptide having one of 10 fully defined sequences given in the (or comprising 40 sequential nucleotides chosen from any of the nucleic acids, its complement or sequences hybridising to it. Also included are a complement or comprising the nucleic acid operably linked to transcription regulatory element, a cell comprising the vector and a single-stranded probe comprising the nucleic acid. The nucleic acids are chosen from 3654 disclosed sequences encoding 3654 disclosed proteins. The nucleic acids is useful for diagnosing pathological conditions resulting from E. faccium bacterial infection (e.g. urinary tract infection, bacteraemia, endocarditis, wounds and abdominal-pelvic infection) and for screening drugs such as agonists and antagonists. The nucleic acid is useful for recombinant production of Candida albicans and very expectations are successful for recombinant production of Candida albicans and very experience of the nucleic acid is useful for recombinant production of Candida albicans and very experience of the nucleic acid is useful for recombinant production of candida albicans and very experience of the nucleic acid is useful for productions.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      and vaccines containing the nucleic acid are useful for preventing or
treating Enterococcus faecium infections. The present sequence represents
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DQIQGGQQLDLSTLNIN-VTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQ 135
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  222 DEIGSGORLDPSTFEINQIHLGEOKVYRGEEGIOOFLODPPSATFNFSVTDNYIEITIPK 281
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Collagen binding Ace protein, antibacterial, immunostimulant, vaccine, extracellular matrix-binding protein, lamin binding; bacterial infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             18 TITSGNKSTNVTVHKSEAG-TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIK 76
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     2;
                  The invention relates to an isolated nucleic acid derived from
                                                                                                                                                                                                                                                                                                                                                                                                                                           43.8%; Score 491; DB 7; Length 731; 48.5%; Pred. No. 2.6e-32; Live 35; Mismatches 64; Indels
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GM, Singh KV,
                                                                                                                                                                                                                                                                                                                                            treating Enterococcus faecium infections. one if the disclosed E. faecium proteins.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Owens RT, Hoo
Weinstock GM,
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(TEXA ) UNIV TEXAS MEDICAL SCHOOL
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TVKGELKVLKQDKDTK 211
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342 TVRGELKIFKYINDTE 357
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Collagen binding Ace protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      99US-0133334P.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Rich RL, Kriekemeyer B,
Nallapareddy SR, Qin X,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WPI; 2000-687639/67.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                L Similarity
95; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                       Sequence 731 AA;
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9
                                                           New collagen-binding protein from Enterococcus, useful e.g. in protective vaccines, for diagnosis and treatment of Enterococcal infections and for screening for compounds that inhibit collagen binding by enterococci.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              IKDOIQGGQGLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIP 134
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               135 QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
                                                                                                                                                                                                                                                        The present sequence is collagen binding Ace protein from Enterococcus faecalis, where Ace is adhesion of collagen from enterococcal bacteria. This protein is an extracellular matrix-binding protein, which can bind with collagens such as collagen type I and type IV and with lamins. Ace protein can be used to inhibit attachment of enterococci to collagen and to treat or prevent enterococcal infections, specifically as a vaccine
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          New cross-reactive monoclonal antibody that binds to a peptide such as the A domain of the ACE protein from Enterococcus faecali, or CAN19 from Staphylococcus aureus, useful for treating or preventing infections
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              18 TITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDITHVRWFLNINNEKSYVSKDIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        cross-reactive monoclonal antibody; ACE; Enterococcus faecalis; antibacterial; veterinary; antibody therapy; vaccine; infection;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Enterococcus faecalis collagen binding ACE protein SEQ ID NO:1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  15.9%; Score 178; DB 3; Length 458; 28.3%; Pred. No. 2.1e-06; ive 33; Mismatches 88; Indels 1
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05-MAR-2002; 2002US-0361347P.
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N-PSDB; ADD90649.
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Matches 54; Conserv
N-PSDB; AAC83828
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Nallapareddy SR,
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                                                      The present invention describes a cross-reactive monoclonal antibody (I) that binds to a peptide selected from the A domain of the ACE protein from Enterococus aureus. Also described: (I) an isolated antisera containing (I); (2) a diagnostic kit comprising (I) and means for detecting binding by (I); (3) a pharmaceutical composition for treating or preventing a bacterial infection comprising (I), and a vehicle, carrier or excipient; (4) a method for treating or preventing aborterial infection comprising (I), and a vehicle, infection of enterococal, or staphylococcal infection by administering (I) to a human or animal patient; and (5) a method for inducing an immunogenic amount of an isolated protein from B. faecalis selected an immunogenic and an isolated protein from B. faecalis selected crown ACE40 and ACE19 protein. (I) has antibacterial and veterinary activities, and can be used in antibody therapy, and in vaccines. The antibody and pharmaceutical composition are useful for treating or preventing infection from staphylococcal, streptococcal and enterococcal according and animal. The present sequence represents the Enterococcus faecalis collagen binding ACE protein, which is used in the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      IKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIP 134
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             180 IADRQGSGQQLNKESFTFDIVNDKETKY---ISLAEFEQQGYGKIDFV--TDNDFNLRFY 234
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Collagen binding Ace protein, antibacterial, immunostimulant, vaccine, extracellular matrix-binding protein, lamin binding, bacterial infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    18 TITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDIT 74
                                                                                                                                                                                                                                                                                                                                                                                                                                      16; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                 ; Score 178; DB 7; Length 458;
; Pred. No. 2.1e-06;
33; Mismatches 88; Indels
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                                                                                                                                                                                                                                                                                                                                          exemplification of the present invention.
                             Claim 8; SEQ ID NO 1; 49pp; English.
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(TEXA ) UNIV TEXAS MEDICAL SCHOOL
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Best Local Similarity 28.3%
Matches 54; Conservative
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caused by the organisms.
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                                                                                                                                                                                                                                                                                                                                                                        Sequence 458 AA;
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New collagen-binding protein from Enterococcus, useful e.g. in protective vaccines, for diagnosis and treatment of Enterococcal infections and for screening for compounds that inhibit collagen binding by enterococci.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       64 KY---ISLAEFEQQGYGKIDFV--TDNDFNLRFYRDKARFTSFIVRYTSTITEAGQHQAT 118
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 The present invention relates to collagen binding Ace protein from Enterococcus faccalis, where Ace is adhesion of collagen from enterococcus bacteria (see AAC83828 and AAB37667). Ace protein is an extracellular matrix-binding protein, which can bind with collagens such as collagen type 1 and type IV and with lamins. Ace protein can be used to inhibit attachment of enterococci to collagen and to treat or prevent enterococcal infections, specifically as a vaccine. The present sequence is the collagen binding domain of Ace protein
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Murray BE;
I KV, Duh
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Owens RT, Hook M, Murra
Weinstock GM, Singh KV,
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(TEXA ) UNIV TEXAS A & M SYSTEM.
(UYPA-) UNIV PAVIA.
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25-APR-2000; 2000US-0199370P.
15-AUG-2000; 2000US-025402P.
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Best Local Similarity 30.29
Matches 45; Conservative
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    Kriekemeyer B,
dy SR, Qin X,
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                                                                                                                         WPI; 2000-687639/67.
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The invention relates to an antibody which recognises a collagen-binding region including CNA19 of CNA protein from Staphylococcus aureus. This antibody is cross-reactive to collagen binding region of both S. aureus and S. epidermidis. It is useful for preventing or treating S. aureus or S. epidermidis infection in human or animal, and for displacing S. aureus or S. epidermidis bound to collagen. Antibody of the invention is useful for interfering with, modulating, and inhibiting the binding interactions between Staphylococcal bacteria and collagen, for detecting the presence of Staphylococcal bacteria or Staphylococcal collagen or binding proteins, to diagnose Staphylococcal infection, as research tools, for
               Novel isolated antibody which recognizes collagen-binding peptide such as CNA19 peptide from Staphylococcus aureus, useful for preventing or treating Staphylococcus aureus or epidermidis infection.
                                                                                                                                                                                                                                                                                                                    development of vaccine for passive immunisation against Staphylococcal infections, and in production facilities or laboratories to isolate additional quantities of collagen-binding proteins. It is also useful for treating medical instruments in order to reduce or eliminate the possibility of them becoming infected or further spreading the infection. The present sequence is Entercoccus faccalis collagen adhesion (ACE)19 protein. This sequence is homologous to S. aureus CNA19 protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           134
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 -----INYKTKIT--NEQQKEFVNNSQAWYQEHGKE 173
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  IKDOIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              18 TITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        50;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                13.7%; Score 154; DB 4; Length 168; 25.8%; Pred. No. 5.5e-05; tive 31; Mismatches 63; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 SdrG; surface protein; infection; antibacterial; vaccine.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           60 IADROGSGOQINKESFTFDIVNDKETKY---ISLABFEQ----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Staphylococcus epidermidis SdrG TR2 protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Ä.
                                                                                             Example 2; Fig 2A; 107pp; English
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABM79017 standard; protein; 316
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (INHI-) INHIBITEX INC.
(TEXA ) UNIV TEXAS A & M SYSTEM.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ATNEKNISQ-VKNV 168
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Staphylococcus epidermidis
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The present sequence comprises the protein sequence of the TR2 protein (amino acids 273-577) of the SdrG surface protein of coagulase-negative Staphylococcus epidermidis. A claimed antibody recognises a protein selected from SdrG NINZN3, NZN3 and TR2. The antibody may be a monoclonal antibody, including a chimeric, murine, humanized, human or single chain monoclonal antibody, which prevents a coagulase-negative staphylococcal infection in a human or animal by inhibiting binding of staphylococcus bacteria to fibrinogen. Such antibodies can be used to treat or prevent staphylococcal infections in cluding nosocomial coagulase-negative staphylococcal infections in low birth weight infants. A claimed vaccine comprises the isolated SdrG NINZN3, NZN3 or TR2 protein
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             161 VEYQKPNENRTANLQSMF-----TNIDTKNHTVEQTIYINPLRYSAKETNVNISGNGD 213
                                                                                                                                                                                             New antibody recognizing a Staphylococcus epidermidis protein comprising SdrG NIN2N3, SdrG N2N3 or SdrGTR2 useful for preparing a composition for treating or preventing a coagulase-negative Staphylococcal infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1 MRGSHHHHHHGSDD-----KVA--TITSGNKSTN--VTVHKSEAGTSSVFY---YKT 45
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 MRGSHHHHHHGSEQGSNVNHLIKVTDQSITEGYDDSDGIIKAHDAE----NLIYDVTFEV 56
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           46 GDMLPEDTTHVRWFLNINNEKSYVSKDIT-----IKDQIQGGQQLDLST---LNINVT
HOOK M;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   75;
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Patel
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                12.6%; Score 141.5; DB 7; 26.5%; Pred. No. 0.0014; ive 32; Mismatches 79;
Domanski P,
MG;
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Hall A,
Bowden N
                                                                                                                                                                                                                                                                                                                                         Claim 27; Page 26; 78pp; English
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Hutchins JT,
Vernachio J,
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                                                                                                  WPI; 2003-722324/68
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                                                                                                                                 N-PSDB; ACF80626
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Sequence 316 AA;
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Patti JM,
Robbins J,
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ABB53995;
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                                                                                                                                                                                                                                                                             The present sequence comprises the protein sequence of the N2N3 region (amino acids 273-597) of the Sdrc surface protein of coagulase-negative Staphylococcus epidermidis. A claimed antibody recognises a protein selected from Sdrc NINZN3, N2N3 and TR2. The antibody may be a monoclonal antibody, including a chimeric, murine, humanized, human or single chain monoclonal antibody, which prevents a coagulase-negative staphylococcal infection in a human or animal by inhibiting binding of staphylococcus bacteria to fibrinogen. Such antibodies can be used to treat or prevent staphylococcal infections including nosocomial coagulase-negative staphylococcal infections in low birth weight infants. A claimed vaccine comprises the isolated SdrG NINZN3, NZN3 or TR2 protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Fibronectin, pCF33; collagen binding protein; sepsis; infection; microbial surface component regonising adhesive matrix molecule; MSCRAMM; adhesin; vaccine; immunisation; diagnosis; therapy; mastitis.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               55 IYDVTFEVDDKVKSGDT-----MTVNIDKNTVPSDLTDSFAIPKIKD--NSGEIIATGTY 107
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              90 --LNINVTGTHSNY---YSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQGYGSYNSFS 144
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ---KT-ITVEYQKPNENRTANLQSMF-----TNIDTKNHTVEQTIYINPLRYSAKETN 211
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      83
                                                                                                                                                                  New antibody recognizing a Staphylococcus epidermidis protein comprising SdrG N1N2N3, SdrG N2N3 or SdrGTR2 useful for preparing a composition for treating or preventing a coagulase-negative Staphylococcal infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ------DDKVATITSGNKSTN--VTVHKSEAGTSSV 40
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNTNKQITYTFTDYVDKYENIKAHLKLTSYIDKSKVPNNNTK--LDVEYKTALSSVN---
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        145 INYKTKITNEQQKEFVN---NSQAWYQEHGKEEVNGKSFNHTVH------
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                                                          Hook
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                                                        Patel P,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11.7%; Score 131.5; DB 7;
25.3%; Pred. No. 0.011;
live 32; Mismatches 78;
                                                        Domanski P,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Firbonectin-binding MSCRAMM derivative pCF33.
                                                      Hall A, Do
Bowden MG;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   -NINANAGIEGTVKGELKVLK 205
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAW31555 standard; protein; 345 AA
                                                                                                                                                                                                                                          Claim 27; Page 25; 78pp; English.
(INHI-) INHIBITEX INC.
(TEXA ) UNIV TEXAS A & M SYSTEM.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        212 VNISGNGDEGSTIIDDSTIIK
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (revised)
(first entry)
                                                      Hutchins JT,
Vernachio J,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1 MRGSHHHHHHGS----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        66; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (revised)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Staphylococcus aureus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Local Similarity
                                                                                                             WPI; 2003-722324/68.
                                                                                                                               N-PSDB; ACF80625
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequence 343 AA;
                                                    Patti JM, F
Robbins J,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            27-AUG-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            25-MAR-2003
21-MAY-1998
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAW31555;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      41
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
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microbial surface component regonising addesive matrix molecule (MSCRAMM) derivative pCF33 plus a vector-derived N-terminal peptide. The invention relates to claimed nucleic acid sequences (see AAT93436-18) encoding S. aureus collagen binding protein (CBP) epitopes M17, M31 and M55 (see AAW31552-54) that confer protection against S. aureus infection. CBP protein and antigenic epitopes are contemplated for use in the treatment of pathological infections, especially to prevent bacterial adhesion to collagen. The epitopes are also contemplated for use in the preparation of vaccines and as carrier proteins in vaccine formulations, as well as in the formulation of compositions for the prevention of S. aureus infection. pCF33 and pQD (see AAW31556) were used to raise anti-MSCRAMM polyclonal antibodies used in passive immunisation against bovine masticis. (Updated on 25-MAR-2003 to correct PI field.) (Updated on 27-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      49 LPEDTTHVRWF-----LNINNEKSYVS-KDITIKDQIQGGQQLDLSTLNINVTGTHSN 100
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      58 VPNSAVKGDTFKITVPKELNLNGVTSTAKVPPIMAGDQVLANGVIDSDG---NVIYTFTD 114
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     YYSGQSAITDFEKA---FPGSKITVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQK 157
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Antibody that interacts with collagen binding domain of Staphylococcal cna gene product - useful to prevent bacterial sepsis in animal infected with Staphylococcus aureus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        1 MRGSHHHHHHGSMVAADAPAAGTDITNQLTNVTVGIDSGTTVYPHQAGYVKLNY---GFS 57
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1 MRGSHHHHHHGS-----DDKVATITSGNKSTNV-----TVHKSEAGTSSVFYYKTGDM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   This protein comprises Staphylococcus aureus fibronectin-binding
                                                                                                                                                                                                                                                                                                                                                                                                                Symersky J;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 2; Length 345;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11.2%; Score 125.5; DB 2; Length 30.1%; Pred. No. 0.035; tive 20; Mismatches 73; Indels
/note= "vector pQE30-derived peptide"
                                                                                                                                                                                                                                                                                                                                                                                                                Sthanam N,
                                                                                                                                                                                                                                                                                                                                                                                                                House-Pompeo K,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Disclosure; Page 91; 143pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABB53995 standard; protein; 336 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Lactococcus lactis protein icaC.
                                                                                                                                                                                                                                                                                                             (TEXA ) UNIV TEXAS A & M SYSTEM (UABR-) UAB RES FOUND.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AUG-2003 to correct OS field.)
                                                                                                                                                                                     97WO-US008210.
                                                                                                                                                                                                                                                    96US-0017678P.
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(first entry)
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                                                                                                                                                                                                                                                                                                                                                                                                                Patti JM,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WPI; 1998-008801/01.
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167 KFYNLS 172
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   158 EFVNNS 163
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
Matches 56; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Sequence 345 AA;
                                                                                                                                                                                                                                                    16-MAY-1996;
                                                              WO9743314-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    29-AUG-2003
16-MAY-2002
                                                                                                                          20-NOV-1997
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             8X4X44X8
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11;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          24 KSTNVTVHKSEAGTSSV------FYYKTGDMLPEDTTHVRWFLNINNEKSYVSK 71
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             58; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          New nucleotide sequence useful in the identification or Lactococcus lactis and related species.
                                    Biosynthesis; biodegradation; lactic bacterium; yogurt; cheese.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.6%; Score 119; DB 5; Length 336; 25.2%; Pred. No. 0.12; tive 28; Mismatches 68; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sorokine A, Renault P, Ehrlich SD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       (INRG ) INRA INST NAT RECH AGRONOMIQUE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Claim 6; SEQ ID NO 697; 2504pp; French.
                                                                                                                                                                                                                                                                                                                11-APR-2000; 2000FR-00004630
                                                                                                                                                                                                                                                                                                                                                                                     11-APR-2000; 2000FR-00004630
                                                                                                 Lactococcus lactis; IL1403
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity 25.2<sup>3</sup>
Matches 52; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WPI; 2002-043418/06.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Sequence 336 AA;
                                                                                                                                                                      FR2807446-A1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Bolotine A,
d
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125 TKNTIDVTIPQGYGSYNSFSINYKTKITN---EQQKEFVNNSQAWYQEHGKEEVNGKSFN 181

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Search completed: July 27, 2004, 10:08:19 Job time : 57 secs

Sequence 2, Appli Sequence 2, Appli Sequence 2, Appli Sequence 67, Appl Sequence 19, Appl Sequence 19, Appl Sequence 3, Appli Sequence 3, Appli Sequence 4, Appli Sequence 14, Appl Sequence 14, Appl Sequence 14, Appl Sequence 14, Appl Sequence 195, Appl

```
Sequence 4, Application US/08856253
; Sequence 4, Application US/08856253
; Patent No. 6288214
; GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: Patti, Joseph M.
APPLICANT: Chause-Pompeo, Karen
APPLICANT: Sthanam, Narashan
APPLICANT: Symersky, Jindrich
TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS
TITLE OF INVENTION: AND METHODS OF USE
NUMBER OF SEQUENCE: 8
CORRESPENDENCE ADDRESS:
ADDRESSE: Annold, White & Durkee
STRRET: P.O. Box 4433
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                CUNTRY: U.S.

ZITT: HOUSEON

ZITT: HOUSEON

ZIP: 77210

COMPUTER: U.S.

ZIP: 77210

COMPUTER: ISP PROPY disk

COMPUTER: IBM PC COMPATIBLE

COMPUTER: IBM PC COMPATIBLE

COMPUTER: PATENTIN Release #1.0, Version #1.30

SOFTWARE: PATENTIN Release #1.0, Version #1.30

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/856,253

FILING APPLICATION 514

PRIOR APPLICATION NUMBER: US 60/017,678

FILING APPLICATION WHERE: US 60/017,678

FILING DATE: LO-MAY-1996

ATTORNEY/AGENT INFORMATION:

NAME: Kitchell, Barbara S.

REGISTRATION NUMBER: 33,928

REFERENCE/DOCKET NUMBER: TAMK:193

TELEPHONE: (512) 418-3000

TELEPHONE: (512) 418-300

TELEPHAX: (512) 418-300
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match 100.0%; Score 1122; DB 3; Best Local Similarity 100.0%; Pred. No. 4.7e-94; Matches 211; Conservative 0; Mismatches 0;
US-08-469-880-2
US-08-728-470-2
US-08-719-641-2
US-08-719-641-2
US-09-206-942-67
US-09-206-942-67
US-09-669-974-19
US-09-669-974-19
US-09-669-974-19
US-09-669-974-19
US-09-797-862-13
US-09-797-862-13
US-08-737-716-14
US-09-543-681A-5491
US-09-543-681A-5491
US-09-543-681A-5491
US-09-563-419-195
US-09-563-419-195
                                                                                                                                                                                                                                                                                  ALIGNMENTS
    ; TOPOLOGY: linear
US-08-856-253-4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STRANDEDNESS:
                                                                                                                                                                                                                                                                                                                              RESULT 1
US-08-856-253-4
    Sequence 10, Appli
Sequence 8, Appli
Sequence 5151, Ap
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Sequence 2, Appli
Sequence 2, Appli
Sequence 3745, Ap
Sequence 7, Appli
Sequence 464, App
Sequence 462, App
Sequence 462, App
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Sequence 30, Appl
Sequence 9, Appli
Sequence 9, Appli
Sequence 9, Appli
Sequence 49, Appli
Sequence 10, Appli
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Sequence 462, App
Sequence 466, App
Sequence 6612, Ap
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                                                                                                                  (without alignments)
573.320 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 2, Sequence 2, Sequence 2,
                                                                                                                                                                                   1 MRGSHHHHHHGSDDKVATIT......GIEGTVKGELKVLKQDKDTK 211
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 10,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Description
                                                                                                  July 27, 2004, 10:07:21 ; Search time 19 Seconds
                                                                                                                                                                                                                                                                                                                                                                                                               Issued Patents AA:*

(cgn2_6/ptodata/2/iaa/5A_COMB.pep:*
(cgn2_6/ptodata/2/iaa/5B_COMB.pep:*
(cgn2_6/ptodata/2/iaa/6A_COMB.pep:*
(cgn2_6/ptodata/2/iaa/6B_COMB.pep:*
(cgn2_6/ptodata/2/iaa/PCTUS_COMB.pep:*
(cgn2_6/ptodata/2/iaa/PCTUS_COMB.pep:*
(cgn2_6/ptodata/2/iaa/pcTuS_COMB.pep:*
                 GenCore version 5.1.6
Copyright (c) 1993 - 2004 Compugen Ltd.
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US-08-856-253-2
US-08-856-253-3
US-08-956-253-7
US-08-956-253-7
US-09-071-035-464
US-09-071-035-466
US-09-071-035-466
US-09-134-000C-6612
US-09-134-000C-6612
US-09-134-000C-6612
US-09-134-000C-6612
US-09-134-000C-6612
US-09-134-000C-6612
US-09-134-00C-6612
US-08-728-470-9
US-08-728-470-9
US-08-728-470-10
US-08-728-470-10
US-08-728-470-10
US-08-728-470-10
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     US-08-038-682-2
US-08-302-832-2
US-08-530-198-2
                                                                                                                                                                                                                                                         389414 segs, 51625971 residues
                                                                                                                                                                                                                                                                                                                                                       Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                       - protein search, using sw model
                                                                                                                                                                                                              BLOSUM62
Gapop 10.0 , Gapext 0.5
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Maximum DB seq length: 2000000000
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                                                                                                                                                        US-09-813-820-4
1122
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Match Length
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125.5
114.5
1114.5
1114.5
1114.5
1114.5
1113.5
1110.5
1100.5
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Perfect score:
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1048
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104.5
104.5
103.5
103.5
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Gaps

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Length 211; Indels

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APPLICANT: GUSS, Bengt
APPLICANT: GUSS, Bengt
APPLICANT: HOOK, Magnus
APPLICANT: JINDBERG, Martin
APPLICANT: LINDBERG, Martin
APPLICANT: SWITALSKI, Lech
APPLICANT: SWITALSKI, SECH
APPLICANT: SWITALSKI, SCH
APPLICANT: SWIT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC Compatible
OPERATING SYSTEM: PC-DOS/WS-DOS
SOFTWARE: PATENTION DATA:
APPLICATION NUMBER: US/08/447,031A
FILING DATE: 22-MAY-1995
CLASSIFICATION: 435
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADDRESSEE: Burns, Doane, Swecker & Mathis STREET: P.O. Box 1404
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 07/861,804
FILING DATE: 21-AUG-1992
PRIOR APPLICATION DATA:
APPLICATION NUMBER: WO PCT/SE91/00707
FILING DATE: 22-OCT-1991
PRIOR APPLICATION NUMBER: SE 9003374-7
FILING DATE: 22-OCT-1990
ATPONEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          186 NINANAGIEGTVKGELKVLKQDKDTK 211
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Sequence 2, Application US/08447031A Patent No. 5851794
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 REGISTRATION NUMBER: 39,300
REFERENCE/DOCKET NUMBER: 01.
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        McGowan, Malcolm K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TELEFAX: (703) 836-2021
INFORMATION FOR SEQ ID NO: SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       STATE: Virginia
COUNTRY: United States
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CORRESPONDENCE ADDRESS:
ADDRESSEE: Burns, Do
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CITY: Alexandria STATE: Virginia
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        rypE: amino acid
ropoLoGY: linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ZIP: 22313-1404
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12
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                                                                                                                                                61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKI 120
                                                                                                                                                                                   61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAPPGSKI 120
                                                                                                                                                                                                                                                                      121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSF 180
                                                                1 MRGSHHHHHHGSDDKVAIITSGNKSINVTVHKSEAGTSSVFYYKTGDMLPEDTIHVRWFL 60
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       GENERAL INFORMATION:
APPLICANT: Hook, Magnus
APPLICANT: Patti, Joseph M.
APPLICANT: Sthanam, Narayana
TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS
TITLE OF INVENTION: AND METHODS OF USE
NUMBER OF SEQUENCES: 8
CORRESPONDENCE ADDRESS:
ADDRESSER: ARDIG, White & Durkee
STREET: P.O. Box 4433
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 3; Length 512;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    STATE: P.O. BOX 4433
CITY: PLOS CONTRY: U.S.
COMPUTER TEXAS
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
OPERATING SYSTEM: PC-DOS/NS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/856,253
FLING DATE: COncurrently Herewith
CLASSIFICATION: 514
PRIOR APPLICATION NUMBER: US 60/017,678
FILING DATE: 16-MAY-1996
ATTORNEY/AGRYT INFORMATION:
NAME: Kitchell, Barbara S.
REGISTRATION NUMBER: 33,228
REFERENCE/DOCKET NUMBER: TAMK:193
TELECOMMUNICATION INFORMATION:
TELEPHONE: (512) 414-3500
TELEFAX: (512) 414-3501
TELEFAX: (512) 414-3517
TELEFAX: (512) 414-3517
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2.1e-87;
0;
                                                                                                                                                                                                                                                                                                                                                                                                181 NHTVHNINANAGIEGTVKGELKVLKQDKDTK 211
                                                                                                                                                                                                                                                                                                                                                                                                                                   0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1 MRGSHHHHHHG-----
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity 64.7
Matches 211; Conservative
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US-08-856-253-6
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US-08-856-253-6
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121 LTQTNTSDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNE 180
                                                                                                                                                                                                                                                                                                                                                          181 KSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNT 240
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     KNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVH 185
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    241 KNTIDVTIPQGYGYGYNSFSINYKTKITNEQQKEFVNNSQAMYQEHGKEEVNGKSFNHTVH 300
61 PTSGTVKIEGYSKTVPLTVKGEQVGQAVITPDGATITFNDKVEKLSDVSGFAEFEVQGRN 120
                                                                                                       12 -----SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNE
                                                                                                                                                                                                                                                                                                      KSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNT
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APPLICANT: Lynn A Doucette-Stamm and David Bush
TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO
ENTEROCOCCUS FAECIUM FOR DIAGNOSTICS AND THERAPEUTICS
                                                                                                                             162 TVTSGDKTATVNVTKPASGSSSVFYYKTGDMLPEDTKHIRWFLNINNGTYVEQPVKIS 221
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 18 TITSGNKSTNVTVHKSEAG-TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIK 76
                                                                   1 MRGSHHHHHHGS-----ITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFL
                                            MRGSHHHHHHGSDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFL
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Gaps
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48.5%; Pred. No. 3.8e-36;
tive 35; Mismatches 64; Indels
                                                                                                                                                                                                                        121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQA 165
                                                                                                                                                                                                                                                    115 TVDNTXNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQA 159
Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               NUMBER OF SEQUENCES: 7310
CORRESPONDENCE ADDRESS:
ADDRESSE: GRNOME THEAPEUTICS CORPORATION
STREET: 100 Beaver Street
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0
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CURRENT APPLICATION DATA:
SOFTWARE: ASCII
CURRENT APPLICATION DATA:
PRIOR APPLICATION DATA:
PRIOR APPLICATION DATA:
PRIOR APPLICATION DATA:
RILING DATE: 14 May 1998
FILING DATE: 14 May 1998
APPLICATION NUMBER: 60/651571
FILING DATE: July 2, 1997
ATTORNEY/AGENT INFORMATION:
0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           NAME/KEY: misc feature
LOCATION: (B) LOCATION 1...731
SEQUENCE DESCRIPTION: SEQ ID NO: 6999:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            NAME: Ariniello, Pamela Deneke
REGISTRATION NUMBER: 40,489
REFERENCE/DOCKET NUMBER: GTC-012
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ORGANISM: Enterococcus faecium
                                                                                                                                                                                                                                                                                                                                                                      Sequence 6999, Application US/09107532A Patent No. 6583275 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                COMPUTER READABLE FORM:
MEDIUM TYPE: CD/ROM ISO9660
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TELECOMMUNICATION INFORMATION:
TELEPHONE: (781)893-5007
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               LENGTH: 731 amino acids TYPE: amino acid
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INFORMATION FOR SEQ ID NO: 6999:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STATE: Massachusetts
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MOLECULE TYPE: protein
HYPOTHETICAL: YES
ORIGINAL SOURCE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity 48.5%
Matches 95; Conservative
  159; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          CITY: Waltham
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ZIP: 02354
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US-09-107-532A-6999
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  Matches
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                                                                                                                                                                                                                                                                                                                                   144 SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSK 203
                                                                                                                                                                                                                                                                                       204 DITIKDÓIQGGQULDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDV 263
                                                                                                                                                                                                                                             72 DITIKDQIQGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDV 131
                                                                                                                                                         71
                                                                                                                                                       12 SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSK
                                                                                                             Gaps
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                                                              Query Match 93.4%; Score 1048; DB 2; Length 1183; Best Local Similarity 100:0%; Pred. No. 2.6e-86; Matches 200; Conservative 0; Mismatches 0; Indels 0
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Pred. No. 8.9e-68;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          GENERAL INFORMATION:
APPLICANT: Hook, Magnus
APPLICANT: Patti, Joseph M.
APPLICANT: House-Pompen
APPLICANT: Sthanam, Narayana
APPLICANT: Symersky, Jindrich
TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS
TITLE OF INVENTION: AND METHODS OF USE
NUMBER OF SEQUENCES: 8
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SOFTWARE PATENTIN Release #1.0, Version #1.30 CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/08/65.253
FILING DATE: Concurrently Herewith CLASSIFICATION NUMBER: US 60/017,678
PRIOR APPLICATION NUMBER: US 60/017,678
FILING DATE: 16-MAY-1996
ATTONNEY/AGBRY INPORMATION:
NAME: Kitchell, Barbara S.
REGISTRATION NUMBER: 33,928
REGISTRATION NUMBER: 33,928
REGISTRATION NUMBER: TAWK:193
TELECOMMULICATION INFORMATION:
TELECOMMULICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5: Arnold, White & Durkee P.O. Box 4433
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   COMPUTER: IBM PC COMPATIBLE OPERATING SYSTEM: PC-DOS/MS-DOS
                                                                                                                                                                                                                                                                                                                                                                                                                           192 GIEGTVKGELKVLKQDKDTK 211
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 324 GIEGTVKGELKVLKODKDTK 343
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Patent No. 6288214
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TELEPHONE: (512) 418-3000
TELEFAX: (512) 474-7577
INFORMATION FOR SEQ ID NO: 2:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        74.0%;
96.4%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      : 159 amino acids amino acids
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COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy
MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ; TOPOLOGY: linear
US-08-856-253-2
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Best Local Similarity
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STREET: P.O. Bo
CITY: Houston
STATE: Texas
COUNTRY: U.S.
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US-08-447-031A-2
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STRANDEDNESS
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                                                                                                                                                                                                                                                                                                                                    GENERAL INFORMATION:
APPLICANT: Lynn Doucette-Stamm et al
TITLE OF INVENTION:
NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO
TITLE OF INVENTION: ENTEROCOCCUS FAECALIS FOR DIAGNOSTICS AND THERAPEUTICS
FILE REFERENCE: 032796-032
CURRENT APPLICATION NUMBER: US/09/134,000C
CURRENT FILING DATE: 1999-08-13
PRIOR FILING DATE: 1997-08-15
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DQIQGGQQLDLSTLNIN-VTGTHSNYYSGQSAITDFEXAFPGSKITVDNTKNTIDVTIPQ 135
                                   222 DEIOSGORLDPSTFEINQIHLGEOKVYRGEEGIQOFLQDFPSATFNFSVTDNYIEITIFK 281
                                                                              136 GYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAGIEG 195
                                                                                                   282 NFVNLKKIMVSYKIIIENPEQINFENHSEAWFKEFNKPAVDGESFNHTVKNISASGGVNG 341
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        75 IKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIP 134
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       135 QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              18 TITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDIT 74
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      88; Indels 16; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             GENERAL INFORMATION:
APPLICANT: Hook, Magnus
APPLICANT: Hotok, Magnus
APPLICANT: House-Pompeo, Karen
APPLICANT: Sthanam, Narayana
APPLICANT: Sthanam, Narayana
APPLICANT: Symersky, Jindrich
TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS
TITLE OF INVENTION: AND METHODS OF USE
NUMBER OF SEQUENCES: 8
CORRESPONDENCE ADDRESS:
ADDRESSES: AROLD, White & Durkee
STREET: P.O. Box 4433
CITY: Houseon
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
15.9%; Score 178; DB 4; Length 540;
Best Local Similarity 28.3%; Pred. No. 5.7e-08;
Matches 54; Conservative 33; Mismatches 88; Indels
                                                                                                                                                                                                                                                                                               Sequence 3745, Application US/09134000C Patent No. 6617156
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Sequence 7, Application US/08856253 Patent No. 6288214
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ORGANISM: Enterococcus faecalis US-09-134-000C-3745
                                                                                                                                                           196 TVKGELKVLKQDKDTK 211
                                                                                                                                                                                                  342 TVRGELKIFKYINDTE 357
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    NUMBER OF SEQ ID NOS: 6812
SOFTWARE: Patentin version 3.1
SEQ ID NO 3745
LENGTH: 540
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                323 VEGEASGNQNV 333
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US-08-856-253-7
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Matches
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Sequence 464, Application US/09071035
Sequence 464, Application US/09071035
Patent No. 6448043
GENERAL INFORMATION:
APPLICANT: Gil H. Choi
TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides
NUMBER OF SEQUENCES: 496
CORRESPONDENCE ADDRESS:
ADDRESSEE: Human Genome Sciences, Inc.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      49 LPEDTTHVRWF-----LNINNEKSYVS-KDITIKDQIQGGQQLDLSTLNINVTGTHSN 100
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        58 VPNSAVKGDTFKITVPKELNLNGVTSTAKVPPIMAGDQVLANGVIDSDG---NVIYTFTD 114
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             101 YYSGQSAITDFEKA---FPGSKITVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQK 157
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1 MRGSHHHHHHGS----DDKVATITSGNKSTNV-----TVHKSEAGTSSVFYYKTGDM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 3; Length 345;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Indels
                   WEDION TYPE: PLOPPY disk
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPRAING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/056,253
FILING DATE: Concurrently Herewith
CLASSIFICATION 514
PRIOR APPLICATION NUMBER: US 60/017,678
FILING DATE: 16-MAY-1996
ATTONNEY/AGBRT INPORMATION:
NAME: Kitchell, Barbara S.
REGISTRATION NUMBER: 33,928
REGISTRATION NUMBER: 33,928
REGISTRATION NUMBER: TAMK:193
TELECOMMUNICATION INPORMATION:
TELEPHONE: (512) 418-3000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ZIP: 20850
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage
COMPUTER: HP Vectra 486/33
OPERATING SYSTEM: MSDOS Version 6.2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       :e 125.5,
d. No. 0.0017;
f. hea 73;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11.2%; Score 125.5;
30.1%; Pred. No. 0.00
tive 20; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                SOFTWARE: ASCII Text
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/071,035
                                                                                                                                                                                                                                                                                                                                                                                                                                                              TELEPHONE: (512) 418-3000
TELEFAX: (512) 474-7577
INFORMATION FOR SEO ID NO: 7:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              LENGTH: 345 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          56; Conservative
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       amino acid
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STATE: Maryland
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RESULT 10
US-09-071-035-462
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Sequence 458, Application US/09071035
Patent No. 6448043
GENERAL INCPORMATION:
GENERAL INCOMMATION:
TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides
NUMBER OF SEQUENCES: 496
CORRESPONDENCE ADDRESS:
ADDRESSEE: Human Genome Sciences, Inc.
STREET: 9410 Key West Avenue
CITY: Rockville
STATE: Maryland
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      71 KDITIKDQIQGGQQLDLSTLNINVTGTHSNYXSGQSAITDFEKAFPGSKITVDNTKN-TI 129
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        : : | : | : | : | : | : | : | : | : : | : : | : : | : : | : : | : : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | 
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     130 DVTIPQGYGSYN-----SPSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   12 SDDKVATITSGNKSTNVTVHKS-EAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVS 70
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB 4; Length 688;
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10.2%; Score 114.5; 1
Best Local Similarity 25.5%; Pred. No. 0.04;
Matches 54; Conservative 36; Mismatches
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CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/071,035
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ATTORNEY/AGENT INFORMATION:
NAME: A. AnderB Brookes
REGISTRATION NUMBER: 36,373
REFERENCE/DOCKET NUMBER: PB369P2
                                                                   FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: A. Anders BXCOOKES
REGISTRATION NUMBER: 36,373
REFERENCE/DOCKET NUMBER: PB36;
TELEPHONE: (301) 309-8504
TELEPHONE: (301) 309-8512
INFORMATION FOR SEQ ID NO: 464:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       688 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ; TOPOLOGY: linear;
; MOLECULE TYPE: protein
US-09-071-035-464
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   single
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COMPUTER READABLE FORM:
MEDIUM TYPE: Diskett
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APPLICATION NUMBER:
FILING DATE:
PRIOR APPLICATION DATA: APPLICATION NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TYPE: amino acid
STRANDEDNESS: si
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          FILING DATE:
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GENERAL INFORMATION:
APPLICANT: Gil H. Choi
TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides
NUMBER OF SEQUENCES: 496
CORRESPONDENCE ADDRESS:
ADDRESSEE: Human Genome Sciences, Inc.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           71 KDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKN-TI 129
                                                                                                                                                                                                                                                                                                                                                                                     908 SYDNTASYINQGSSRDVTGKVSIQHGGESV--KKGGEYHKDDPDHVYWHVMINGAQS-VL 964
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
                                                                                                                                                                                                                                                                                                                                                      12 SDDKVATITSGNKSTNVTVHKS-EAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVS 70
                                                                                                                                                                                                                                                                                                            29;
                                                                                                                                                                                                                                                                DB 4; Length 2032;
                                                                                                                                                                                                                                                                                                            93; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage COMPUTER: HP Vectra 486/33
OPERATING SYSTEM: MSDOS version 6.2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DDNGDVVVDIDHSGGHATGTKGKIQLKKTAMD 1104
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              178 KSFNHTVHNINANAGIEGTVKGELKVLKQDKD 209
                                                                                                                                                                                                                                                                10.2%; Score 114.5; I
25.5%; Pred. No. 0.2;
tive 36; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              STREET: Human Genome Sciences, Inc. STREET: 9410 Key West Avenue CITY: Rockville STATE: Maryland COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   SOFTWARE: ASCII Text
CURENT APPLICATION DATA:
APPLICATION NUMBER: US/09/071,035
FILING DATE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              REGISTRATION NUMBER: 36,373
REFERENCE DOCKET NUMBER: P8369P2
TELECOMMUNICATION INFORMATION:
TELEPHONE: (301) 309-8504
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Sequence 462, Application US/09071035
Patent No. 6448043
             TELEPHONE: (301) 309-8504
TELEFAX: (301) 309-8512
INFORMATION FOR SEQ ID NO: (458:
SEQUENCE CHARACTERISTICS:
LENGTH: 2032 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TELEFAX: (301) 309-8512
INFORMATION FOR SEQ ID NO: 462:
SEQUENCE CHARACTERISTICS:
LENGTH: 2032 amino acids
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: A. Anders Brookes
REGISTRATION NUMBER: 36,
                                                                                                                                                                                                                                                           Query Match
Best Local Similarity 25.5%
Matches 54; Conservative
                                                                                                                                                     single
                                                                                                                                                                     TOPOLOGY: linear
MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER:
                                                                                                                             TYPE: amino acid
STRANDEDNESS: si
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GENERAL INFORMATION:
APPLICANT: Lynn Doucette-Stamm et al.
TITLE OF INVENTION:
TITLE OF INVENTION:
TITLE OF INVENTION:
FILE REFERENCE: 032796-032
CURRENT APPLICATION NUMBER: US/09/134,000C
CURRENT FILING DATE: 1999-08-13
FILE REFERENCE: 1999-08-13
FILE REFERENCE: 1999-08-13
FILE REFERENCE: 1999-08-13
FILE REFERENCE: 1999-08-13
FRIOR FILING DATE: 1997-08-15
                                            71 KDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKN-TI 129
                                                                                                                                    130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 71 KDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKN-TI 129
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
  908 SYDNTASYTNQGSSRDVTGKVSIQHGGESV--KKGGEYHKDDPDHVYWHVMINGAQS-VL 964
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   12 SDDKVATITSGNKSTNVTVHKS-EAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 2054;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         93;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1095 DDNGDVVVDIDHSGGHATGTKGKIQLKKTAMD 1126
                                                                                                                                                                                                                                                                    1073 DDNGDVVVDİDHSGGHATGTRGKIQLKKTAMD 1104
                                                                                                                                                                                                                          178 KSFNHTVHNINANAGIEGTVKGELKVLKQDKD 209
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          178 KSFNHTVHNINANAGIEGTVKGELKVLKQDKD 209
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.2%; Score 114.5; DE
25.5%; Pred. No. 0.21;
ive 36; Mismatches
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Patent No. 6077692
GENERAL INFORMATION:
APPLICANT: RUBEN, STEVEN M.
APPLICANT: JIMENEZ, PABLO
APPLICANT: APPLICANT: APPLICANT: APPLICANT: APPLICANT: APPLICANT: APPLICANT: APPLICANT: MGNORE, PAUL A.
APPLICANT: MOORE, PAUL A.
APPLICANT: COLEMAN, TIMOTHY A.
APPLICANT: GOLEMAN, TIMOTHY A.
APPLICANT: GOLEMAN, TIMOTHY A.
APPLICANT: GOLEMAN: JOACHIM R.
APPLICANT: DILLON, PATRICK J.
                                                                                                                                                                                                                                                                                                                                                                          Sequence 6612, Application US/09134000C Patent No. 6617156
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TYPE: PRT ORGANISM: Enterococcus faecalis
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               NUMBER OF SEQ ID NOS: 6812
SOFTWARE: PatentIn version 3.1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity 25.5*
Matches 54; Conservative
                                                                                                                                                                                                                                                                                                                                                               US-09-134-000C-6612
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   US-09-134-000C-6612
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US-09-071-035-466
US-09-071-035-466
Sequence 466, Application US/09071035
Patent No. 6448043
GENERAL INFORMATION:
APPLICANT: Gil H. Choi
TITLE OF INFORMIN: Enterococcus faecalis Polynucleotides and Polypeptides
NUMBER OF SEQUENCES: 496
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                           9
                                                                                                                                                                                   908 SYDNTASYTNÇGSSRDVTGKVSIQHGGESV--KKGGEYHKDDPDHVYMHVMINGAQS-VL 964
                                                                                                                                                                                                                                               71 KDITIKDOIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKN-TI 129
                                                                                                                                                                                                                                                                                                                                       130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
                                                                                                                                                         12 SDDKVATITSGNKSTNVTVHKS-EAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVS 70
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   12 SDDKVATITSGNKSTNVTVHKS-EAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVS 70
                                                                                                               29;
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                                                                   Length 2032;
                                                                                                               Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Indels
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COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage
COMPUTER: HP Vectra 486/33
OPERATING SYSTEM: MSDOS version 6.2
SOFTWARE: ASCIT Text
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/071,035
                                                                                                             93;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match 10.2%; Score 114.5; DB 4; Best Local Similarity 25.5%; Pred. No. 0.2; Matches 54; Conservative 36; Mismatches 93;
                                                                 DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                     | : | : : | | | 104
                                                                                                                                                                                                                                                                                                                                                                                                                              178 KSFNHTVHNINANAGIEGTVKGELKVLKQDKD 209
                                                              Query Match
10.2%; Score 114.5; D
Best Local Similarity 25.5%; Pred. No. 0.2;
Matches 54; Conservative 36; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADDRESSEE: Human Genome Sciences, Inc. STREET: 9410 Key West Avenue CITY: Rockville STATE: Maryland
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      PB369P2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION:
APPLICATION UNBER:
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME:
REGISTRATION NUMBER: 36,373
REFERENCE/DOCKET NUMBER: PB365
TELEPHONE: (301) 309-8504
TELEPHONE: (301) 309-8512
INFORMATION FOR SEQ ID NO: 466:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             2032 amino acids
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, MOLECULE TYPE: protein US-09-071-035-462
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           TYPE: amino acid
STRANDEDNESS: sir
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       linear
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58 LFSFTKYFLK--IEKNGKVSGTKKENCPYSILEITSVEIGVVAVKAINSNYY---LAMNK 112
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            111 FEKAFPGSKITVD-NTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQE 169
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1 MRGSHHHHHHGSDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFL 60
                                                                                                                       US-09-248-998-30

IS Sequence 30, Application US/09248998

Patent No. 6598799

GENERAL INFORMATION:
APPLICANT: Gimenez, Pablo
APPLICANT: Rampy, Mark A.
APPLICANT: Russell, Deborah
APPLICANT: Louie, Arthur
TITLE OF INVENTION: Therapeutic Uses of Keratinocyte Growth Factor-2
FILE REPERENCE: 1488-1066002
CURRENT APPLICATION NUMBER: US/09/248,998
CURRENT PILING DATE: 39-99-02-12
FARLIER APPLICATION NUMBER: US 60/114,387
EARLIER PILING DATE: 30-DEC-1998
EARLIER PILING DATE: 13-FEB-1998
NUMBER OF SEQ ID NOS: 148
SOFTWARE: Patentin Ver. 2.0
SEQ ID NO 30
LENGTH: 184
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Sequence 9.288651

Fatent No. 528851

GENERAL INFORMATION:

APPLICANT: Barenkamp, Stephen J

TITLE OF INVENTION: High Molecular Weight Surface Proteins

TITLE OF INVENTION: of No. 5928651-Typeable Haemophilus

TITLE OF INVENTION: of No. 5928651-Typeable Haemophilus

NUMBER OF SEQUENCES: 10

CORRESPONDENCE ADDRESS: 10

CORRESPET: 2001 Jefferson Davis Hwy., 1203 Crystal Plaza

STREET: Shoemaker and Mattare, Ltd.

STREET: Bldg. 1

CITY: Alington

STATE: Virginia

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     47;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 184;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            113 KGKLYGSKEFNNDCKLKERIE---ENGYNTYASFN---
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity 23.49
Matches 45; Conservative
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148 NGROMYVALNGK 159
       148 NGROMYVALNGK 159
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US-09-248-998-30
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US-08-728-470-9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         111 FEKAFPGSKITVD-NTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQE 169
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ----WOH 147
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               STREET: STERNE, KESSLER, GOLDSTEIN & FOX, P.L.L.C. STREET: 1100 NEW YORK AVE, NW, SUITE 600 STATE: DC COUNTRY: USA 210.
                                                                                                                                                                                                                                                                                                                           COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
COMPTOTE: IBM PC compatible
COMPTOTE: IBM PC compatible
COMPTOTE: IBM PC compatible
SOFTWARE: Batentin Release #1.0, Version #1.30
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/023,082A
FLING DATE: 13-FEB-1995
CLASSIFICATION NUMBER: WS 08/461,195
FILING DATE: 14-FEB-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 00/023,852
FILING DATE: 13-AUG-1996
FILING DATE: 13-AUG-1996
PRIOR APPLICATION NUMBER: US 00/039,045
FILING DATE: 28-FEB-1997
PRIOR APPLICATION NUMBER: US 00/039,045
FILING DATE: 13-AUG-1997
PRIOR APPLICATION NUMBER: US 00/039,045
FILING DATE: 13-AUG-1997
PRIOR APPLICATION NUMBER: US 08/910,875
FILING DATE: 13-AUG-1997
PRIOR APPLICATION NUMBER: US 08/910,875
FILING DATE: 13-AUG-1997
PRIOR APPLICATION NUMBER: US 08/910,875
FILING DATE: 13-AUG-1997
ATTORNEY/AGENT INFORMATION:
NUMBER: STEFFEFE ERIC K.
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APPLICANT: GENTZ, REINER L.
TITLE OF INVENTION: KERATINOCYTE GROWTH FACTOR-2
NUMBER OF SEQUENCES: 148
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      65;
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10.1%; Score 113.5; DB
Best Local Similarity 23.4%; Pred. No. 0.0088;
Matches 45; Conservative 35; Mismatches 6
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                NAME: STEPFE, ERIC K.
REGISTRATION NUMBER: 36,688
REGISTRATION NUMBER: 1488.0360008/EKS
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-371-2600
INFORMATION FOR SEQ ID NO: 30:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             HGKE---EVNGK 178
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MOLECULE TYPE: protein
US-09-023-082A-30
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        121 TVDNTKNTIDVTIPQGYGSY------NSFSINYKTKITNEQQKEFV 160
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      12 SDDKVATITSGNKST----NVTVHKSEA----GTSSVPYYKTGDMLPEDTTHVRWFLN 61
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 :|| :| || :| || 852 DNSTGLTISAKDVTVVNNVTSHKTINISABAAGNVTTKEGTTINATTGSVEVTAQNGTIK 910
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          161 NNSQAWYQEHGKEEVNGKSFNHTVHNINANAG----IEGT----VKGELKVLKQDKDTK 211
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
9.8%; Score 110.5; DB 2; Length 1338;
Best Local Similarity 25.5%; Pred. No. 0.26;
Matches 61; Conservative 28; Mismatches 89; Indels 61; Gaps
APPLICATION NUMBER: US/08/728,470
FILING DATE:
CLASSIFICATION: 424
PRIOR APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION DATA:
APPLICATION NUMBER: US PCT/US93/02166
FILING DATE: 16-MAR-1993
FILING DATE: 16-MAR-1993
ATORNEY/AGENT INFORMATION:
BEGISTRATION NUMBER: 22,651
ATORNEY/AGENT INFORMATION:
TELEPHONE: (703) 415-0813
TELECOMUNICATION INFORMATION:
TELEPHONE: (703) 415-0813
INFORMATION FOR SEQ ID NO: 9:
SEQUENCE CHARACTERISTICS:
LENGTH: 1338 amino acids
TYPE: amino acids
TYPE: amino acids
TYPE: amino acid
STRANDEDNESS: single
170POLOGY: linear
US-08-728-470-9
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Search completed: July 27, 2004, 10:10:29 Job time: 20 secs

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July 27, 2004, 10:06:36; Search time 16 Seconds (without alignments) 1268:526 Million cell updates/sec
                                                                                                                                                                                            US-09-813-820-4
1122
1 MRGSHHHHHHGSDDKVATIT......GIEGTVKGELKVLKQDKDTK 211
GenCore version 5.1.6
Copyright (c) 1993 - 2004 Compugen Ltd.
                                                                              - protein search, using sw model
                                                                                                                                                                                                                                                                             BLOSUM62
Gapop 10.0 , Gapext 0.5
                                                                                                                                                                                                Title:
Perfect score:
                                                                                                                                                                                                                                                                               Scoring table:
                                                                              OM protein
                                                                                                                                                                                                                                        Sequence:
                                                                                                                    Run on:
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283366 Total number of hits satisfying chosen parameters:

283366 segs, 96191526 residues

Searched:

seq length: 0 seq length: 200000000 Minimum DB Maximum DB Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

PIR 78:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

disconctin-bindin collagen adhesin [hypothetical prote peptidoglycan anch high-molecular-wei hypothetical prote wall associated pr probable surface f hypothetical prote fibronectin-bindin autolysin [importe neurexin III beta finger protein MSN hypothetical prote lysozyme 2 (EC 3.2 ABC-type transport hypothetical prote cell surface antig surface anchored p DNA topoisomerase hypothetical prote. hypothetical prote hypothetical prote toxin-like outer m Clost DNA topoisomerase collagen adhesin neurotoxin -cell surface Description SUMMARIES A42404 A48620 B42798 B96710 A98021 A41396 A4 E86706 A42296 C48399 E90065 C71953 C89921 B71704 AC1458 S73898 B64596 T06029 % Query Match Length DB 911 131 1131 1100 1000 99.5 99.5 98.5 98 Š

C;Species: Staphylococcus arreus C;Species: Staphylococcus arreus C;Date: 07-Apr-1994 #sequence_revision 18-Nov-1994 #text_change 03-Mar-1995 C;Accession: A46820 R;Patti, J.M.; Boles, J.O.; Hook, M. Biochemistry 32, 11428-11435, 1993 A;Title: Identification and biochemical characterization of the ligand binding domain of A;Reference number: A48620, MUID:94032261; PMID:8218209 A;Contents: FDA 574 A;Accession: A48620 A;Status: preliminary; not compared with conceptual translation

toxin-like outer m

A48620 adhesin - Staphylococcus aureus (fragment)

RESULT 2

DNA-directed RNA p	transcription fact	hypothetical prote	hypothetical prote	opacity protein op	probable exported	sensory transducti	hypothetical prote	hypothetical prote	cellulase (EC 3.2.	ferrichrome-bindin	lipoprotein a [imp	hypothetical prote	rhoptry protein -	apolipoprotein B-1	аврагаgine-rich pr
T28156	T13350	G86643	G71609	S16619	AF0169	871629	F90073	AE1710	A44808	H97200	B90578	C89824	T28677	JH0101	S14535
~	N	~	N	7	N	7	N	N	7	N	N	N	7	7	~
2910	1037	1983	2500	261	622	2150	2271	452	463	471	578	953	2269	784	699
8.2	8.5	8.2	8.2	8.2	8.2	8.2	8.2	8.1	8.1	8.1	8.1	8.1	8.1	8.1	8.0
92.5	92	92	92	91.5	91.5	91.5	91.5	91	91	91	91	91	91	90.5	90
30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

ALIGNMENTS

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C;Species: Streptococcus pneumoniae
C;Species: Streptococcus pneumoniae
C;Species: Streptococcus pneumoniae
C;Date: 22-Oct-2001 #sequence_revision 22-Oct-2001 #text_change 22-Oct-2001
C;Accession: Ag0021
R;Hoskins, J.A.; Alborn Jr., W.; Arnold, J.; Blaszczak, L.; Burgett, S.; DeHoff, B.S.; E. e, R.; LeBlanc, D.J.; Lee, L.N.; Lefkowitz, E.J.; Lu, J.; Matsushima, P.; McAhren, S.; M.; P.; Sun, P.M.; Winkler, M.E.
J; Bacteriol. 183, 5709-5717, 2001
A;Authors: Yang, Y.; Young-Bellido, M.; Zhao, G.; Zook, C.; Baltz, R.H.; Jaskunas, S.R.; A;Reference number: A97872; MUID:21429245; PMID:11544234
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C;Species: Listeria monocytogenes
C;Species: Listeria monocytogenes
C;Species: Listeria monocytogenes
C;Accession: AH1396
R;Glaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloecker
J; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H.
D; Jones, L.M.; Katst, U.
Science 294, 849-852, 2001
A;Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Ma.
A;Authors: Comparative genomics of Listeria species.
A;Title: Comparative genomics of Listeria species.
A;Reference number: AB1077; MUID:21537279; PMID:11679669
                                           : | | | : | | | : | | : | | : | | : | | : | | : | | : | | EDTRVVVDWKDGGTTEVNIGPATGPDKDEVLYKWGWVDENDSTLIHWQVRINYAKENIQK 130
                                                                                                                                                                                A; Cross-references: GB: AE007317; PIDN: AAK99997.1; PID: 915458827; GSPDB: GN00174 C; Genetics:
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                                                                                                                                                                                                                                                                                                                                                  -----YENRG--ELTGENIE 221
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                                                                                                                                            DITIKDQIQGGQQL----DLSTLNINVTGTHSNY----YSGQSAITDFEKAFPGSKITVDN
   --FYYKTGDMLPEDTTHVRWFLNINNEKSYVSK
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                                                                                                                                                                                                                                                                                                                                                                                                                      182 -----HTVHN-INANAGIEGTVKGE 200
                                                                                                                                                                                                                                                                                                                                                                                                                                                             222 KQVVEVHTPNNGGNGNASMKLSISGE 247
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   : | : | | 108 SENPLE-LNNSSFW 210
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       KSTNVTVHKSEAGTSSV
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les 49; Conserv
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A;Molecule type: DNA
A;Residues: 1-542 <KUR>
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Cipacies: Lactococcus lactis subcoccus
Cipacies: Lactococcus
Cipacies: Lact
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 124 NTKNTIDVTIPQ----GYGSY----NSFSINYKTKITNEQQKEFVNNSQAMY-QEHGKE 173
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            266 FSGQTVSINIPEKIQYDNGSYQELNESEMQIHLYAKI----KKEVLENSSIEYVRNESKV 321
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                collagen adhesin [imported] - Lactococcus lactis subsp. lactis (strain IL1403) C;Species: Lactococcus lactis subsp. lactis 
C;Date: 23-Mar_2001 #sequence_revision 23-Mar-2001 #text_change 03-Aug-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          73
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     fibronectin-binding protein - Streptococcus "equisimilis"
C;Species: Streptococcus "equisimilis"
C;Date: 06-Jan-1995 #sequence_revision 06-Jan-1995 #text_change 15-Oct-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    C,Accession: 842798
R,Signaes, C.

R,Signaes, C.

A,Accession: 842798
A,Accession: 842798
A,Scatus: preliminary
A,Molecule Cype: DNA
A,Residues: 1-1092 < SIGS
C,Kesidues: EMBL:Z29088; NID:g436135; PIDN:CAA82330.1; PID:g436136
C,Keywords: fibronectin binding
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       19 ITSGNKSTNVTVHKSEAGTSS-VFYYKTGDMLPEDTTH---VRWFLNIN-NEKSYVSKDI
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                                                                                                                                                                                                                    Gaps
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                                                                                                                                                                                                                ;
0
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                                                                                                                                            Length 37;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          11.7%; Score 131; DB 2; Length 10
26.5%; Pred. No. 0.13;
tive 31; Mismatches 88; Indels
                                                                    A; Note: sequence extracted from NCBI backbone (NCBIP:138726)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         EVNGKSF----NHT---VHNINANAGIEGTVKGELKVLK 205
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 || |::: | | ::| | | EVRGENWPIDPNSTSAYVQILRQGGWAKGTVRGEVRILK 360
                                                                                                                                                                                                                                                                                                                     1 ITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTH 37
                                                                                                                                        Query Match 17.3%; Score 194; DB 2; L
Best Local Similarity 100.0%; Pred. No. 5.2e-08;
Matches 37; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                 19 ITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTH
   nucleic acid; protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity 26.5%
Matches 58; Conservative
                                       <PAT>
Molecule type:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      174
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96

Gaps

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C;Species: Clostridium acetobutylicum
C;Date: 14-Sep-2001 #sequence_revision 14-Sep-2001 #text_change 14-Sep-2001
C;Accession: D97203
R;Nolling, J.; Barteon, G.; Omelchenko, M.V.; Markarova, K.S.; Zeng, Q.; Gibson, R.; Lee, S;Nolling, J.; Barteon, G.; Moonin, B.V.; Smith, D.R.
J; Bally, M.J.; Bennett, G.N.; Koonin, B.V.; Smith, D.R.
A;Title: Genome Sequence and Comparative Analysis of the Solvent-Producing Bacterium Clc. A;Reference number: A96900; MUID:21359325; PMID:21359325
A;Accession: D97203
A;Accession: D97203
A;Accession: D97203
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AG1542

wall associated protein precursor (LPXTG motif) homolog lin0879 [imported] - Listeria in C; Species: Listeria innocua
C; Species: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 27-Nov-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Cjaccesion: AGIS42
R;Glaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, P.; Berche, P.; Bloecker, D.; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H.D.; Jones, L.M.; Karet, U.S.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H.D.; Jones, L.M.; Karet, U.S.; Kuhn, M.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Mahathors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Mahathors: Kreft, J.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland, A; Reference number: AB1077; MUID:21537279; PMID:11679669
A; Status: preliminary
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    A;Residues: 1-463 <GLA>
A;Residues: 1-463 <GLA>
A;Cross-references: GB:AL592022; PIDN:CAC96111.1; PID:g16413329; GSPDB:GN00178
A;Experimental source: strain Clip11262
C;Genetics: A;Gene: 1in0879
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SO YYNSNDGKEVERSKIITHPINNKSKPFIVERVKIGTDTKIITYEDDYSCRLIEPMVDKKVK 109
                                                                                                                                                                                                                                                                                                                             A,Cross-references: GB:AE001437; PIDN:AAK80415.1; PID:g15025480; GSPDB:GN00168
A,Experimental source: Clostridium acetobutylicum ATCC824
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  99 SNYYSGQSAITDFEKAFPGSKITVD----NTKNTIDVTI----PQGYGSYNSFSINYKTK 150
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      164 GDNFDKN----TEAQKVYSFMKKIVDTKLVKTHMHIEVSFYFVKSQYYNKTNPQAIAYRYN 220
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 76 KDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQ 135
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              212 TDTLGTGQNL-LGNATIKYRDANKKVLS----TNVQ-----PIALDSNRN-FELTI-- 256
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            47 DMLPEDTTHVRW-----FLNINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTH
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               19 ITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           53;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 266;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         151 ITNEQQKEFVN----NSQAWYQEHG--KEEVNGKSFNHTVHNINAN 190
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   221 -RSMMDSEWLNKNENTTKNYVDHSFTLRSEDGLSYNQDINEIMKN 264
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          78; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              9 HHGSDD------KVATITSGNKSTNVTVHKSEAGTSSVFYY-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         83;
                                                                                                                                                                                                                                                                                                                                                                                                                                                         8.9%; Score 99.5; DB 2;
22.2%; Pred. No. 4.1;
iive 39; Mismatches 83;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 2;
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; Pred. No. 8.3;
33; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity 24.3%;
Matches 50; Conservative 3:
                                                                                                                                                                                                                                                                                                                                                                                                                                                 Ouery Match
Best Local Similarity 22.2%
Matches 50; Conservative
                                                                                                                                                                                                                                                                                                          A; Residues: 1-266 <KUR>
                                                                                                                                                                                                                                                                             A; Molecule type: DNA
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high-molecular-weight surface-exposed protein HWW1 - Haemophilus influenzae
CiSpeciaes: Haemophilus influenzae
CiPecession: A43855
RiBarenkamp, 8.J.; Leininger. E.
Infect. Immun. 60, 1302-1313, 1992
A;Title: Cloning, expression, and DNA sequence analysis of genes encoding nontypeable fidetella pertussis.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             A;Reference number: A43855; MUID:92192797; PMID:1548058
A;Accession: A43855
A;Actatus: preliminary
A;Molecule type: DNA
A;Residues: 1-1536 <BAR>
A;Residues: 1-1536 <BAR>
A;Roses-references: GB:U08876; GB:M84616; NID:g475770; PIDN:AAA20527.1; PID:g475771
A;Note: sequence inconsistent with the nucleotide translation
A;Note: sequence extracted from NCBI backbone (NCBIN:89235, NCBIP:89239)
       A;Accession: AH1396
A;Status: preliminary
A;Moleculs type: DNA
A;Rossiduss: 1-1530 <GLA>
A;Cross-references: GB:NC_003210; PIDN:CAD00654.1; PID:gl6412064; GSPDB:GN00177
B;Experimental source: strain EGD-e
C;Genetics:
A;Gene: lmo2576
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                                                                                                                                                                                                                                                                                                        10;
                                                                                                                                                                                                                                                                                                                                                                                            | | | : : | | | : | | | : | | 665 EWKLIFNDSSKLISKP-TIEDSLNSGQTFIQDSIEIHKIDLSATPQVGELIPPENYDVTF 665
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                719
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       943 NSKNLSITINSSSTYRTI-----ISGNITNKNGDLNITNEGSDTEMQIGGDVSQKEGNL 996
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         61 NINNEKSYVSKDITIKDQIQGQQLDLSTLNINVT-GTHSNYYSGQSAITDFEK----AF 115
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           PGSKITVDNTKNTIDVTIPQGYGSYNSFSINYK----TKITNEQQKEFVNNSQAWYQEHG 171
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           97 THSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQGYGSYNSFSI-----NY 147
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          9
                                                                                                                                                                                                                                                                                                                                                                                                                                                            RWFLNINNEKSYVSKDITIKDQIQGGQ------QLDLSTL-----NINVTG 96
                                                                                                                                                                                                                                                                                                                                                       -- VFYYKTGDMLPEDTTHV
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           | : | | | | : | : | | | | : | | TKKENGNEQMLIT-FKKPLI-HPVEVTYKTKPVGITKP----LYKNKAVISDGEEVLADY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       13 DDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDM-------LPEDTTHVRWFL
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D97203
hypothetical protein CAC2461 [imported] - Clostridium acetobutylicum
                                                                                                                                                                                                                                                                                                        64;
                                                                                                                                                                                                                                                Length 1530;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 2; Length 1536;
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                                                                                                                                                                                                                                                                                                        Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  73;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     72;
                                                                                                                                                                                                                                                   Query Match 8.9%; Score 100; DB 2; Best Local Similarity 22.4%; Pred. No. 34; Matches 49; Conservative 33; Mismatches 73
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     33; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   8.9%; Score 100;
25.2%; Pred. No. 3
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                                                                                                                                                                                                                                                                                                                                                          17 ATITSG-NKSTNVTVHKSEAGTSS-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Conservative
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Matches 51; Conserv
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75

Gaps

1104 RIGKIYAĞRDILTFNKSNAĞĞKSEIINRĞTINVKNKLSYDS-DVS 1146 Qy		22491. 22491.	A;Residues: 1-1117 <lin> A;Residues: 1-1117 <lin> A;Cross-references: EMBL:22151; NID:9288970; PIDN:CAA80122.1; PID:9581643 A;Note: the authors translated the initiation codon TTG for residue 1 as Leu C;Genetics: A;Gene: fnbB A;Start codon: TTG C;Keywords: fibronectin binding F;1-27/Domain: signal sequence #status predicted <ngt> F;28-1117/Product: fibronectin-binding protein #status predicted <nat></nat></ngt></lin></lin>	Query Match 8.7%; Score 98; DB 2; Length 1117; Best Local Similarity 23.3%; Pred. No. 32; Matches 48; Gaps 11; Matches 51; Conservative 35; Mismatches 85; Indels 48; Gaps 11; Qy 27 NVTVHKSEAGTSSVFXKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIKDQIQ 80 I 1 1 1 1 1 1 Db 157 NINYNAGGSRTDSV-YAKSGVLDDFDSTGRIVWYFNFNAALNPI-KDEDVEFMTTDNLP 214	Qy 81 GGQQLDLSTLNINVTGTH-SNYYSGQSAITDFEKAFPGSKITVDNTK 126	QY 127 NTIDVIPQGYGSYNSPSINYKTKITNEQOKEFVNNSQAWYQEH 170	RESULT 13 C89874 autolysin [imported] - Staphylococcus aureus (strain N315) C;Species: Staphylococcus aureus C;Species: Staphylococcus aureus C;Species: 10-May-2001 #sequence_revision 10-May-2001 #text_change 22-Oct-2001 C;Accession: C89874 R;Kuroda, M.; Ohta, T.; Uchiyama, I.; Baba, T.; Yuzawa, H.; Kobayashi, I.; Cui, L.; ma, A.; Mizutani-Ui, Y.; Kobayashi, N.; Sawano, T.; Inoue, K.; Kaito, C.; Sekimizu,	Lancet 357, 1225-1240, 2001 A/Title: Whole genome sequencing of meticillin-resistant Stapylococcus aureus. A/Reference number: A89758; MUID:21311952; PMID:11418146
OY 136 GYGSYNSFSINYKTKITNBOOKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINAN 190	bril protein NMA1200 [imported] - Neisseria meningitidis (s a meningitidis #sequence_revision 05-May-2000 #text_change 02-Feb-2001	Accession: AB1888 A; Accession: AB175; MulD:2022556; PMID:10761919 A; Reference number: AB1775; MUID:2022556; PMID:10761919 A; Reference number: AB1775; MUID:2022556; PMID:10761919 A; Reference number: AB1775; MUID:2022556; PMID:10761919 A; Residues: preliminary A; Molecule type: DNA A; Residues: 1-592 < PAR> A; Residues: 1-592 < PAR> A; Coss-references: GB:AL162755; GB:AL157959; NID:g7379742; PIDN:CAB84461.1; PID A; Genetics: NAA1200	Query Match 8.9%; Score 99.5; DB 2; Length 592; Best Local Similarity 21.6%; Pred. No. 11; Additional States of Sta	60 LNINNEKSYVSKDITIKDQIQG	Qy 180 FNHTVHNINANAGIEGTVK 198 : Db 301 TDEGEGLVTAKEVIDAVNK 319	TESULT 11 T31105 T931105 T931105 T931105 T931105 T931105 C.Species: Haemophilus ducreyi C.Species: Haemophilus ducreyi C.Species: Asemophilus ducreyi C.Species: 22-Oct-1999 #sequence_revision 22-Oct-1999 #text_change 22-Oct-1999 C.Accession: T31105 TS.F. Latimer, J.L.; Cope, L.D.; Hansen, E.J. T. Racteriol 180. 6013-6022, 1998	Arritle: Haemophilus ducrey; secretes a filamentous hemagglutinin-like protein. A; Reference number: 220984; MUID:99030326; PMID:9811662 A; Accession: T31105 A; Accession: T31105 A; Molecule type: DNA A; Residues: 1-4919 < WAR> A; Residues: 1-4919 < WAR> A; Cross-references: EMBL:AF057696; NID:g3929021; PID:g3929023; PIDN:AAC79761.1 C; Genetics: A; Gene: 1spA2	Query Match 8.8%; Score 98.5; DB 2; Length 4919; Best Local Similarity 24.8%; Pred. No. 1.9e+02; Matches 53; Conservative 34; Mismatches 70; Indels 57; Gaps 12;

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meurexin III beta precursor (clone pB794-7) - bovine
C;Species: Bos primigenius taurus (cattle)
C;Date: 12-Apr-1995 #sequence_revision 12-Apr-1995 #text_change 09-Dec-2002
C;Date: 12-Apr-1995 #sequence_revision 12-Apr-1995 #text_change 09-Dec-2002
C;Accession: A5380
R;Ushkaryov, X.A.; Hata, Y.; Ichtchenko, K.; Moomaw, C.; Afendis, S.; Slaughter, C.A.; E. J. Biol. Chem. 269; 11994
A;Title: Conserved domain structure of beta-neurexins. Unusual cleaved signal sequences A;Reference number: A5380
A;Accession: A5380
A;Accession: A5380
A;Accession: A5380
A;Accession: A5380
A;Residues: 1-392 <USH>A
A;Cross-references: GB:L27864
C;Superfamily: neurexin; EGF homology
C;Keywords: alternative splicing
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  93 NVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQGYGSYNSFSIN--YKTK 150
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     151 ITNEQQKEFVNNS-----QAWYQEHGKEEVNGKSFNHTVHNINANAGIEGTVKGEL 201
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     96
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    58 WFLN----INNEKSYVSKDITIKDQI-------QG--GQQLDLSTLNI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       84;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ; Score 96; DB 2;
; Pred. No. 12;
35; Mismatches 74
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              completed: July 27, 2004, 10:09:59
He : 17 secs
193 IEGTVKG---ELKVLKQDK 208
                                           691 QEYTKKGKPKKLKIIKAKK
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity 19.6%;
Matches 47; Conservative 3:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Job time
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R; Fraser, C.M.; Casjens, S.; Huang, W.M.; Sutton, G.G.; Clayton, R.; Lathigra, R.; White Son, D.; Peterson, J.; Kerlavage, A.R.; Quackenbush, J.; Salzberg, S.; Hanson, M.; Vugt, Bob. D.; Peterson, J.; Kerlavage, A.R.; Quackenbush, J.; Salzberg, S.; Hanson, M.; Vugt, Son, D.; Se0-586, 1997
Nature 390, 580-586, 1997
Nature 390, 580-686, 1997
A; Aluthors Smith, H.O.; Venter, J.C.
A; Atthors Smith, H.O.; Venter, J.C.
A; Reference number: A70100; MUID: 98065943; PMID: 9403685
A; Accession: C70203
A; Accession: C70203
A; Accession: C70203
A; Residues: 1-848 «KLE>
A; Residues: 1-848 «KLE>
A; Residues: 1-848 «KLE>
A; Residues: Leferences: Strain B31
C; Superimental source: strain B31
C; Superimental source: strain B31
C; Superfamily: bacterial type I DNA topoisomerase
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               958 SDTAKYSLKAFNEOPFAVVKEQVINGOTWY--YGKLS-NGKLAWIKSTDLAKELIKYNOT 1014
                                                                                                                                                                                                                                                                                                                                   16;
                       A;Residues: 1-1248 «KUR»
A;Residues: 1-1248 «KUR»
A;Cross-references: GB:BA000018; PID:g13700854; PIDN:BAB42150.1; GSPDB:GN00149
A;Experimental source: strain N315
C;Genetics:
A;Gene: atl
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     841
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            842 NVKDLNVQNLGKEVKTTQKYTVNKSNNGLSMVPWGTKNQVILTGNN----IAQGTFNATK 897
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     630
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      113 KAFPGSKI----TVDNTK---NTIDVTIP---------QGYGSY----N 141
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        142 SFSINYKTKITNEQQ----KEFVNNSQAWYQEHGKEEVNGK-----SFNHT 183
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         69 VSK-DITIKDQIQGGQQLDL----STLNINVTGTH------SNTYS 103
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           --FEKAFPGSKITVDNTKNTIDVTIPQG-YGSY----NSFS 144
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   89
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA topoisomerase I (topA) homolog - Lyme disease spirochete
C,Species: Borrelia burgdorferi (Lyme disease spirochete)
C,Date: 13-Feb-1998 #sequence_revision 13-Feb-1998 #text_change 16-Jul-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                          574 GKKGLKDTVMQLEPKIDSSEFRTVIESQKI---ENKNSINYTINIGKYGPYLIFKGHNYS
                                                                                                                                                                                                                                                                                                                                                                                                              KVATITSGNKSTNVTVHKSEAGTSSVP----YYKTGD------MLPEDTTH---VRWF
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
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                                                                                                                                                                                                                                                                                                                                   90;
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                                                                                                                                                                                                                                                    Query Match 8.7%; Score 98; DB 2;
Best Local Similarity 21.6%; Pred. No. 37;
Matches 61; Conservative 39; Mismatches 90
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         8.6%; Score 96.5; DB 22.8%; Pred. No. 29; ative 41; Mismatches
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Best Local Similarity 22.8%;
Matches 59; Conservative
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clostridium fusobacteri pieris bras chlamydia p saccharomyc saccharomyc saccharomyc

P44969 P44584 P45384 P5384 P5331 P53659 Q8rf47 Q99736 P48361 Q926570 P56570 P56570 P38078

candida tro

lactococcus buchnera ap

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      STRAIN=FDA 574;
MEDLINE=94032261; PubMed=8218209;
Patti J.M., Boles J.O., Hoeoek M.;
"Identification and biochemical characterization of the ligand
binding domain of the collagen adhesin from Staphylococcus aureus.";
Biochemistry 32:11428-11435(1993).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               aureus adhesin."

Nat. Struct. Biol. 4:833-838(1997).

-!- FUNCTION: MEDIATES ATTACHMENT OF STAPHYOCOCCAL CELLS TO COLLAGEN-CONTAINING SUBSTRATA.

-!- SUBCELLULAR LOCATION: Attached to the cell wall peptidoglycan by an amide bond (Potential).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   MEDLINE=97475225; PubMed=9334749; Symersky J., Patti J.M., Carson M., House-Pompeo K., Teale M., Moore D., Jin L., Schneider A., DeLucas L.J., Hoeoek M., Narayana S.V.L.; Raryana S.V.L.; Structure of the collagen-binding domain from a Staphylococcus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 STRAIN=FDA 574;
MEDLINE-92165839; PubMed=1311320;
Patti J.M., Jonsson H., Guss B., Switalski L.M., Wiberg K.,
Lindberg M., Hoeoek M.;
Molecular characterization and expression of a gene encoding
Staphylococcus aureus collagen adhesin.";
J. Biol. Chem. 267:4766-4772(1992).
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Lindberg M., Hoeoek M.;
J. Biol. Chem. 269:11672-11672(1994).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Staphylococcus aureus.
Bacteria, Firmicutes, Bacillales, Staphylococcus
NCBI_TaxID=1280;
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Last annotation update)
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IGAO HAEIN
ACMA HAEIN
ACMA LACLC
Y3.7 BUCAL
GUNZ CLOSR
ACOB FUSNN
PRSN PIEBR
PM15 CHLPN
AS10 YEAST
IRSX YEAST
PPZ1 YEAST
VATA CANTR
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15-DEC-1998 (Rel. 37, Last seq
10-OCT-2003 (Rel. 42, Last ann
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P33046

P39046

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Copyright (c) 1993 - 2004 Compugen Ltd.
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Gapop 10.0 , Gapext 0.5
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AMIDE-LINKED TO CELL WALL (POTENTIAL).
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                                                                                                                                                                                                                                                                                                           COLLAGEN ADHESIN.
REMOVED BY SORTASE (POTENTIAL).
COLLAGEN-BINDING.
3 X 187 AA APPROXIMATE TANDEM REPEALYS/PRO-RICH (CELL WALL-SPANNING).
B1.
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                                                                                                                                                                                                                                                                          Peptidoglycan-anchor; Repeat; Signal; 3D-structure
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                                                                     InterPro; IPR008966; Adhes bact.
InterPro; IPR008956; Adhes bact.
InterPro; IPR008454; Cna B.
InterPro; IPR008970; Cna B.
InterPro; IPR008956; Collagen bind.
InterPro; IPR001899; Gram_pos=anchor.
Pfam; Pr05738; Cna B; 7.
Pfam; Pr05738; Cna B; 7.
ITGREAMS; TIGR0167; LPXTG anchor; 1.
TIGREAMS; TIGR0167; LPXTG anchor; 1.
PROSITE; PS50847; GRAM_POS_ANCHORING; FALSE_NEG.
Cell wall; Peptidoglycan-anchor; Repeat; Signal; 31
SIGNAL
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1AMX; 24-JUN-98.
1D20; 27-SEP-00.
1D2P; 27-SEP-00.
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                                                                                                                                                                                                                                                                                                                                                                                               Kruhoeffer M., Cetin Y., Kaempf U., Forssmann W.-G.;
Submitted (AUG-1996) to the EMBL/GenBank/DBM databases.
-!- FUNCTION: RECEPTOR FOR THE B. COLI HAT-STABLE ENTEROTOXIN (E.COLI
ENTEROTOXIN MARKEDLY STIMULATES THE ACCUMULATION OF COMP IN
MAMMALIAN CELLS EXPRESSING GC-C). ALSO ACTIVATED BY THE ENDOGENOUS
                                                                                                                                                          ົວ
                        01-NOV-1997 (Rel. 35, Created)
01-NOV-1997 (Rel. 35, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
Heat-stable enterotoxin receptor precursor (GC-C) (Intestinal
guanylate cyclase) (EC 4.6.1.2) (STA receptor) (Guanylyl cyclase (GCC)
                                                                                                                                                                                                           Cavia porcellus (Guinea pig).
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Hystricognathi; Caviidae; Cavia.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   -1- CATALYTIC ACTIVITY: GTP = 3',5'-cyclic GMP + diphosphate.
-1- SUBCELLULAR LOCATION: Type I membrane protein.
-1- SIMILARITY: Belongs to the adenylyl cyclase class-4/guanylyl cyclase family.
-1- SIMILARITY: Contains 1 protein kinase-like domain.
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PROSITE; PS50125; GUÁNYLATE_CYCLASES_2; 1.
PROSITE; PS5011; PROVERN KITANSE_DOM; 1.
Receptor; Transmembrane; Ğlycoprotein; Lyase; cGMP biosynthesis;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        POTENTIAL.
HEAT-STABLE ENTEROTOXIN RECEPTOR.
EXTRACELLULAR (POTENTIAL).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            9B53F16E05E80EB9 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      CYTOPLASMIC (POTENTIAL).
PROTEIN KINASE-LIKE.
GUANYLATE CYCLASE.
O'LINKED (GLCNAC. . . ) (PO'N-LINKED (GLCNAC. .
PRT; 1076 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        POTENTIAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 InterPro; IPR001828; ANF receptor. InterPro; IPR001054; G_Cyclase. InterPro; IPR000719; Prot kinase. InterPro; IPR000719; Prot kinase. Pfam; PF01094; ANF_receptor; 1. Pfam; PF00211; guanylate Cyc; 1. Pfam; PF00069; pkinase; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PRINTS; PRO0109; TYRKINASE.
ProDom; PD000001; Prot kinase; 1.
SMART; SM00044; CXCc; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               EMBL; Z74734; CAA98989.1; -. HSSP; Q02846; 1AWL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 123119
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Receptor; Transmembrane;
Signal. 1 23
SIGNAL 24 1076
STANDARD;
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457
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957
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                                                                                                                                                                                                                                                                                                                                                SEQUENCE FROM N.A.
TISSUE=Colon;
                                                                                                                                                                                                                                                                                              NCBI_TaxID=10141;
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310
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SEQUENCE
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Score 106; DB 1; Length 1076;

9.48;

Query Match

RESULT 2

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                 10;
                                                              KASNLPFKSFSWNTSYVF--KNG----TESEHCFWYINALEAGVSYFSQVLGFKEMLRGN 232
                                                                                      -----LSTLNINVTGTH-----SNYYSGQSAITDF 111
                                                                                                                                                                                                         334 LFGHMLKTFLRNG-ENTTAHKFAHAFRNLTFEGSTGPVTLDDSGDIDNTMVLLYTSVDTK 392
                                        82
                                                                                                                                                                                                                                                                                      01-0CT-1996 (Rel. 34, Last sequence update)
01-0CT-1996 (Rel. 34, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
Bifunctional autolysin precursor [Includes: N-acetylmuramoyl-L-alanine amidase (EC 3.5.1.28); Mannosyl-glycoprotein endo-beta-N-acetylglucosamidase (EC 3.2.1.96)].
                                      24 KSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLN-INNEKSYVSKDITIKDQIQGG
                                                                                                                                                  ------DYMKNVLVLTLPPGNSTIN-----TSLSKESLQEFSDFALAYLDGIL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       -----YQEHGKEEVNGKSFNHTVHNINANAGI-----EGTVKGELKVLKQDKDTK
                                                                                                                                    112 EKAFPGSKITVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQOKEFVNNSQAW----
                  Gape
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      glycopeptides.
CATALYTIC ACTIVITY: Endohydrolysis of the di-N-acetylchitobiosyl
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        unit in high-mannose involved to the containing the -[Man (GLCNAc)2] Asn-structure. One N-acety1-D-glucosamine residue remains attached to the protein; the rest of the oligosaccharide is ratleased intact.
SUBCELLULAR LOCATION: Secreted.
PTM: UNDERGOES PROTEOLYTIC PROCESSING TO GENERATE THE TWO EXTRACELLULAR LYTIC EMXZMES.
SIMILARITY: IN THE N-TERMINAL SECTION; BELONGS TO THE N-ACETYLMURAMOYL-L-ALANINE AMIDASE PAMILY 2.
SIMILARITY: IN THE C-TERMINAL SECTION; BELONGS TO FAMILY 73 OF GLYCOSYL HYDROLASES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      "A Staphylococcus aureus autolysin that has an N-acetylmuramoyl-L-alanine amidase domain and an endo-beta-N-acetylglucosaminidase domain: cloning, sequence analysis, and characterization."; proc. Natl. Acad. St. U.S.A. 92:285-289(1995)
                 78;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Oshida T., Sugai M., Komatsuzawa H., Hong Y.-M., Suginaka H.,
                 Indels
                                                                                                                                                                                                                                                                                                                                                                                                                             SEQUENCE FROM N.A., AND SEQUENCE OF 205-214 AND 776-792.
                78;
                                                                                                                                                                                                                                                                                                                                                                                         Bacteria, Firmicutes, Bacillales, Staphylococcus
      No. 2.6;
                                                                                                                                                                                                                                                                       1256 AA
19.6%; Pred. ....
                                                                                                                                                                                                                                                                                                                                                                                                                                                    MEDLINE=95116542; PubMed=7816834;
                47; Conservative
                                                                                                                                                                                                                                                                      STANDARD;
                                                                                                                                                                                                                                                                                                                                                                               Staphylococcus aureus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                STRAIN=NCTC 8325-4;
                                                                                      83 QQLD-----
     Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                       NCBI_TaxID=1280;
                                                                                                                                                                                                                                                                                                                                                                                                                                         STRAIN=RN450;
                                                                                                                                                                                                                                                                      STAAU
                                                              179
                                                                                                                                                                                    167
                                                                                                                                                            292
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Tomasz A.
                Matches
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modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ----LNINN--EKSYVSKDIŢIKDQIQGGQQLDLSTLN-INVŢGTHSNYYSGQSAITDFE 112
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              850 NVKDLAVQULGKEVKTTQKYTVNKSNNGLSMVPWGTKNQVLLTGNN----IAQGTFNATK 905
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      -----QGYGSY----N 141
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       906 OVSVGKDVYLYGTINNRTGWVNAKDLTAPTAVKPTTSAAKDYNYTYVIKNGNGYYYVTPN 965
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            142 SFSINYKTKITNEQQ----KEFVNNSQAWYQEHGKEEVNGK-----SFNHT 183
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        790 KIAQVKPNNTGIRASVYEKTAKNGAKYADRIFÝVTKERAHGNETYVLLNNTSHNIPLGWF
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gwinn M.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         15 KVATITSGNKSTNVTVHKSEAGTSSVF----YYKTGD------MLPEDTTH---VRWF
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Praser C.M., Cadjens S., Huang W.M., Sutton G.G., Clayton R.A., Lathigra R., White O., Ketchum K.A., Dodson R., Hickey E.K., Gwinn M Dougherty B. Tomb J.-F., Fleischmann R.D., Richardson D., Peterson J., Kerlavage A.R., Quackenbush J., Salzberg S., Hanson M., van Vugt R., Palmer N., Adams M.D., Gocayne J.D., Weidman J., Utterback T., Watthey L., McDonald L., Artiach P., Bowman C., Garland S., Fujii C., Cotton M.D., Horst K., Roberts K., Hatch B., Smith H.O., Veneer J.C.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          15-DEC-1998 (Rel. 37, Created)
15-DEC-1998 (Rel. 37, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
DNA topoisomerase I (EC 5.99.1.2) (Omega-protein) (Relaxing enzyme)
(Untwisting enzyme) (Swivelase).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Borrelia burgdorferi (Lyme disease spirochete).
Bacteria, Spirochaetes; Spirochaetales; Spirochaetaceae; Borrelia.
                                                                                                                                                                                                                                                                                                                                                                                               BIFUNCTIONAL AUTOLYSIN.
N-ACETYLMURAMOYL-L-ALANINE AMIDASE.
ENDO-BETA-N-ACETYLGLUCOSAMIDASE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  92;
                                                                                                                                                                                                                                                                                                                                                Multifunctional enzyme; Repeat.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 98; DB 1; Length 1256; Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     90; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         137384 MW; 2BB76CAA292FDD20 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        184 VHNINANAGIEGTVKGELKV-------LKQDKDTK
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      113 KAFPGSKI----TVDNTK---NTIDVTIP----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             848 AA
                                                                                                                                                                                                                                                                                                                                                                            POTENTIAL
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STRAIN=ATCC 35210 / B31;
MEDLINE=98065943; PubMed=9403685;
                                                                                                                                                                                InterPro; IPR002502; Amidase_2.
InterPro; IPR002501; Amidase_4.
Pfam; PF01510; Amidase_4.
Pfam; PF01832; Amidase_4; 1.
SMART; SM00644; Ami_2; 1.
SMART; SM00647; LYZ2; 1.
Cell wall; Hydrolase; Signal; Mul
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     39;
                                                                                                                                 EMBL; D17366; BAA04185.1; -. EMBL; L41499; AAA99982.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 8.7%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        61; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                               1256
775
1256
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1256 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
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REPEAT
REPEAT
                                                                                                                                                                                                                                                                                                                                                                      SIGNAL
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             14;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          145 INYKTKITNEQQKE----FVNNSQAWYQEHGKEEVNGKS--FNHTVH-NI----NANAG 192
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               68
                                                -:- FUNCTION: The reaction catalyzed by topoisomerases leads to the conversion of one topological isomer of DNA to another.
-:- CATALYTIC ACTIVITY: ATP-independent breakage of single-stranded DNA, followed by passage and rejoining.
-:- SUBUNIT: Monomer (By similarity)
-:- MISCELLANEOUS: When a topoisomerase transiently breaks a DNA backbone bond, it simultaneously forms a protein-DNA link, in which a tyrosyl oxygen in the enzyme is joined to a DNA phosphorus at one end of the enzyme-severed DNA strand.
-:- SIMILARITY: Belongs to the prokaryotic type I/III topoisomerase
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  19 ITSGNKSTNVTVHKSEAGTSSVFYY-----KTGDMLPEDTTHVRWFLNINNEKSY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 104 GQSAITD-----FEKAFPGSKITVDNTKNTIDVTIPQG-YGSY----NSFS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    8.6%; Score 96.5; DB 1; Length 848; 22.8%; Pred. No. 9.9; ive 41; Mismatches 84; Indels 7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SWARK; SW00436; TUPLDC, SWART; SW00436; TUPLDC, SWART; SW00493; TOPRIM; 1.

TIGRFAMS; TIGRO1051; topA bact; 1.

PROSITE; PS00396; TOPOISOMERASE I PROK; 1.

ISOMERASE; TOPOISOMERASE I DROK; 1.

ISOMERASE; TOPOISOMERASE; DAA-binding; Complete proteome.

ISOMERASE; TOPOISOMERASE; DAA-binding; COMplete proteome.

324 324 324 DNA CLEAVAGE (BY SIMILARITY).
sequence of a Lyme disease spirochaete, Borrelia
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         69 VSK-DITIKDQIQGGQQLDL----STLNINVTGTH---
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       InterPro; IPR005733; DNA_topl_bact.
InterPro; IPR000380; DNA_tpisomrase.
InterPro; IPR003601; DNAtopl_ATP bind.
InterPro; IPR003602; DNAtopl_DNA_bind.
InterPro; IPR006171; Toprim_dom.
InterPro; IPR006154; Toprim_dub.
Fam; PF01131; Topoisom_bac; 1.
Pfam; PF01131; Topoisom_bac; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  193 IEGTVKG---ELKVLKQDK 208
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          691 QEYTKKGKPKKLKIIKAKK 709
                                                                                                                                                                                                                                                                                                                                                                                                                                   EMBL; AE001180; AAC67161.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PRTPISMRASEI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PRINTS; PRO0417; PRTPISMRA
SMART; SM00437; TOP1AC; 1.
SMART; SM00436; TOP1BC; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Local Similarity 22.8%
nes 59; Conservative
                                     Nature 390:580-586(1997).
                                                                                                                                                                                                                                                                                                                                                                                                                                                      PIR; C70203; C70203.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        P06612; 1ECL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           BB0828;
                                                                                                                                                                                                                                                family.
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NX3B_BOVIN
ID NX3B_BOVIN
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Matches
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456 AA

PRT;

STANDARD;

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ALTERNATIVE PRODUCTS:

Event-Alternative splicing; Named isoforms=4;

Event-Alternational isoforms seem to exist. There is a

Comment-Additional isoforms seem to exist.

Combination of two alternatively spliced domains at sites 4 and

combination of two alternatively spliced domains at sites 4 and
                                                                                                                                                                                                                                                                                                                                                  Unusual cleaved signal
                                                                                                                                                                                                                                                                                       MEDLINE=94216308; PubMed=8163501; Ushkaryov Y.A., Hata Y., Ichtchenko K., Moomaw C., Afendis S., Slaughter C.A., Suedhoff T.C.; Staughter C.A., Suedhoff T.C.; Suedhoff T.C.; Suedhoff T.C.; Suedhoff T.C.; Suedhoff T.C.; Suedis structure of beta-neurexins. Unusual cleaved signa sequences in receptor-like neuronal cell-surface proteins."; J. Biol. Chem. 269:11987-11992(1994).
-!- FUNCTION: Neuronal cell surface protein that may be involved in cell recognition and cell adhesion.
-!- SUBUNIT: The cytoplasmic C-terminal region binds to CASK (By similarity). Binds to neuroligins NLGNI, NLGNZ and NLGNI (By
                                                                                                                                    Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi;
ammaila; Butheria, Cetartiodactyla, Ruminantia, Pecora, Bovoidea,
Bovidae, Bovinae, Bos.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  - SIBLIAILLY).
- SUBCELLULAR LOCATION: Type I membrane protein (Potential)
ISOforms 3 and 4 seems to be secreted.
- ALTERNATIVE PRODUCTS:
                                                                                                                                                                                                                                  SEQUENCE FROM N.A. (ISOFORMS 1; 2; 3 AND 4), AND SEQUENCE OF
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             NEUREXIN 3-BETA.
EXTRACELLULAR (POTENTIAL)
POTENTIAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              IsoId=028143-4; Sequence=VSP_003533, VSP_003534;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5, which seem to be used independently. Experii confirmation may be lacking for some isoforms;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PROSITE, PS50025; LAM G DOMAIN, 1.
Signal; Transmembrane; Cell adhesion; Glycoprotein;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Note=Lacks the transmembrane domain, SIMILARITY: Contains 1 laminin G-like domain. SIMILARITY: Belongs to the neurexin family.
                28-FEB-2003 (Rel. 41, Created)
28-FEB-2003 (Rel. 41, Last sequence update)
20-CCT-2003 (Rel. 42, Last annotation update)
Neurexin 3-beta precursor (Neurexin III-beta)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 IsoId=Q28143-3; Sequence=VSP_003534;
Note=Lacks the transmembrane_domain;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Name=2;
IsoId=028143-2; Sequence=VSP_003533;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       IsoId=Q28143-1; Sequence=Displayed;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              HSSP; Q63373; 1C4R.
InterPro; IPR008985; ConAlike lec_gl
InterPro; IPR001791; Laminin G.
InterPro; IPR003585; Neurexin-like.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  EMBL; L27869; AAA19908.1; ALT_INIT.
PIR; A53580; A53580.
PIR; B53580; B53580.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Pfam; PF00054; laminin G; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    EMBL; L27869; AAA19907.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Alternative splicing.
SIGNAL 135
CHAIN 36 456
DOMAIN 36 381
TRANSNEM 382 402
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SMART; SM00294; 4.1m; 1
SMART; SM00282; LamG; 1
                                                                                                                  Bos taurus (Bovine)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              similarity)
                                                                                                                                                                                                                                                      N-TERMINUS.
TISSUE=Brain;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Name=1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DOMAIN
TRANSMEM
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   229 SDINSYSNSISNSNSNSTGNLN----SSYFNSLNIDSMLDDYVSSDLLLNDDDDDTNLSR 284
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               -----STLNINVTGTHSNYYSGQSAIT 109
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               343 TTMNANADSNI-AGNPKNNDATİDNELTQILNEYNMNFNDNLGTSTSGKNKSACPSSFDA 401
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                146 NYKTKITNEQQ-KEFVNNSQ--AWYQEHGKEEVNGKSFNHTVHNINANAG---IEGTVKG 199
                                                                                                                                                                                                                                                                                                                          J. Cell Biol. 161:497-505(2003).

-!- FUNCTION: POSITIVE TRANSCRIPTIONAL FACTOR THAT ACTS AS A COMPONENT OF THE STRESS RESPONSE SYSTEM. RECOGNIZES AND BINDS TO THE STRESS RESPONSE ELEMENT (STRE) WHICH IS INVOLVED IN THE RESPONSE INVOLVED IN THE RESPONSE INVOLVED IN THE REGULATION OF THE CTT1, DDR2, HSP12 GENES.
-!- SUBCELLULAR LOCATION: Cytoplasmic and nuclear.
-!- SIMILARITY: Contains 2 C2H2-type zinc fingers.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       12 SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSK
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         285 RRFSDV-ITNOFPSMTNSRNSISHSLDLWNHPKINPSNRNTNLNIT-TNSTSSSNASPNT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   PROSITE; PS00028; ZINC FINGER C2H2 1; 2.
PROSITE; PS50157; ZINC_FINGER_C2H2_2; 2.
DNA-binding; Nuclear protein; Zinc-finger; Metal-binding; Activator;
                    MEDLINE=96208509; PubMed=8641288;
Martinez-Pastor M.T., Marchler G., Schueller C., Marchler-Bauer A.,
                                                                               "The Saccharomyces cerevisiae zinc finger proteins Msn2p and Msn4p are required for transcriptional induction through the stress response element (STRE).";
EMBO J. 15:2227-2215(1996).
                                                                                                                                                                                                                    MEDLINE=22628141; PubMed=12732613; De Mey J., Goldbeter A.; Jacquet M., Renault G., Lallet S., De Mey J., Goldbeter A.; Soscillatory nucleocytoplasmic shuttling of the general stress response transcriptional activators Msn2 and Msn4 in Saccharomyces
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               90;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                8.6%; Score 96; DB 1; Length 704;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             97; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      77860 MW; EDF6F07446819DF1 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          110 DFEKAFPGSKITVDNTKN---TIDVTIPQGYGSYN------
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (ACIDIC)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ASP-RICH (ACIDIC)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     8.5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             31; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ASP-RICH (ACI C2H2-TYPE 1. C2H2-TYPE 2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        stress; IMP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               72 ----DITIKDQIQG------GQQLDL-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Transcription regulation; Repeat.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          InterPro; IPR007087; Znf C2H2.
Pfam; PF00096; zf-C2H2; Z.
ProDom; PD000003; Znf C2H2; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 EMBL; L08838; AAA34806.1; -.
EMBL; Z48502; CAA88403.1; -.
PIR, 339004, S39004.
HSSP; P08047; ISP1.
GermOnline; 142706; -.
TRANSFAC; T01257; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              GO; GO:0005829; C:cytosol; IDA.
GO; GO:0005634; C:nucleus; IDA.
GO; GO:0006550; P:response to EinterPro; IPR007087; Znf. C2H2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     25.4%;
                                                                                                                                                                              [4]
NUCLEOCYTOPLASMIC SHUTTLING
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              C2H2; 2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               64; Conservative
                                                              Ruis H., Estruch F.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         S0004640; MSN2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                SMART; SM00355; ZnF
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ZN_FING
ZN_FING
SEQUENCE
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  g
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                                                          POLY-THR.
POLY-THR.
POLY-THR.
POLY-ALA.
N-LINKED (GLCNAC. . .) (POTENTIAL).
N-LINKED (GLCNAC. . .) (POTENTIAL).
N-LINKED (GLCNAC. . .) (POTENTIAL).
Missing (in isoform 2 and isoform 4).
MISSING (in isoform 2 and isoform 4).
ANPTERGYREVERSESSITCHAVOGIVAAAALCILI
LIYAMYKYRNRDEGSYQVDETRNYISNSAGSNGTLLKEKPP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     93 NVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKVTIDVTIPQGYGSYNSFSIN--YKTK 150
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             157 SIKEERTPVNDGKYHVVRFTRNGGNATLQVDN----------WPVNEHYPTG 198
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      199 NTDNERFQMVKQKIPFKYNRPVEEWLQEKGRQ------LTIFNTQAQIAIGGKDKGRL 250
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          151 ITNEQQKEFVNNS-----QAWYQEHGKEEVNGKSFNHTVHNINANAGIEGTVKGEL 201
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SSKGGHKKQKNKDKEYYV -> ARSSNAARSLRAALTWTWR
LTYTFTPIIFISCVVHS (in isoform 3 and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5 HHHHHHGSDDKVATITSGNKSTNVTVHKS-----EAGTSSVFYYKTGDMLPEDTTHVR
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Estruch F., Carlson M.; "Two homologous zinc finger genes identified by multicopy suppression in a SNF1 protein kinase mutant of Saccharomyces cerevisiae."; mol. cell. Biol. 13:3872-3881(1993).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ------OG--GOOLDLSTLNI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      84; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE FROM N.A.
STRAIN=S288C / AB972;
MEDLINE=97313268; PubMed=9169972;
MEDLINE=97313268; PubMed=9169972;
Connor R., Churcher C.M., Badcock K., Brown D., Chillingworth T., Jogens R., Dedman K., Devlin K., Gentles S., Hamlin N., Hunt S., Jagels K., Lye G., Moule S., Odell C., Pearson D., Rajandream M.A., Rice P., Skelton J., Walsh S., Whitehead S., Barrell B.G.;
With nucleotide sequence of Saccharomyces cerevisiae chromosome XIII.";
Nature 387:90-93(1997).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           01-FEB-1994 (Rel. 28, Created)
01-FEB-1994 (Rel. 28, Last sequence update)
10-OCT-2003 (Rel. 42, Last annotation update)
Zinc finger protein WNU2 (Multicopy suppressor of SNF1 protein 2)
MSN2 OR YMR037C OR YM9532.02C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Saccharomyces cerevisiae (Baker's yeast).
Eukaryota, Fungi, Ascomycota, Saccharomycotina, Saccharomycetes,
Saccharomycetales, Saccharomycetaceae, Saccharomyces.
                                                                                                                                                                                                                                                                                                                                                                                                                          DB 1; Length 456;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      74; Indels
                                                                                                                                                                                                                                                                                                                                                        /FTId=VSP_003534.
E1B5467089A8CF2C CRC64;
CYTOPLASMIC (POTENTIAL)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      35; Mismatches
                         G-LIKE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    704 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           58 WFLN----INNEKSYVSKDITIKDQI-----
                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity 19.6%; Pred. No.
Matches 47; Conservative 35; Mismatc
                                                                                                                                                                                                                                                                                                                                        isoform 4)
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                                                                                                                                                                                                                                                                                                                                                                                  50244 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                  456 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SEQUENCE FROM N.A. STRAIN=S288c;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         01-FEB-1994
01-FEB-1994
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 MSN2 YEAST
                                                                                                                                                                                                                                                                                                                                                                                SEQUENCE
                                                                                                                                  CARBOHYD
                                                                                                                                                       CARBOHYD
                                                                                                                                                                                                                                             VARSPLIC
                                                                                                                                                                                                 VARSPLIC
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        TRAINMANCE 9790;

A Del Mar Lleo M., Canepari P., Satta G.;

Del Mar Lleo M., Canepari P., Satta G.;

Thermosensitive cell growth mutants of Enterococcus hirae that

"Thermosensitive cell growth are something of the cell marchiol. 139:3099-3117(1993).

"Gen. Mircobiol. 139:3099-3117(1993).

"The cell marchiol. 139:3099-3117(1993).

"The cell marchiol. 139:3099-3117(1993).

"The mutants The Norman are stimulated to divide by

"The cell marchiol. 139:3099-3117(1993).

"The cell marchiol. 139:309-3117(1993).

"The cell marchiol. 13
Dolinger D.L., Daneo-Moore L., Shockman G.D.;
"The second peptidogly/van hydrolase of Streptococcus faecium ATCC 9790
covalently binds penicillin.";
J. Bacteriol. 171:4355-4361(1989).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Enterococcus hirae.
Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DOMAIN: LysM repeats are thought to be involved in peptidoglycan
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-FEB-1995 (Rel. 31, Last sequence update)
Last annotation update)
Muramidase-2.precursor (EC 3.1.17) (1,4-beta-N-
acetylmuramoylhydrolase) (Peptidoglycan hydrolase) (Pg-hydrolase-2)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Chu C.-P., Kariyama R., Daneo-Moore L., Shockman G.D.; "Cloning and sequence analysis of the muramidase-2 gene from
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             SIMILARITY: Belongs to family 73 of glycosyl hydrolases.
                                                                                                                                                                                                                                                                                                                                                                                    666 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SEQUENCE FROM N.A., AND SEQUENCE OF 50-73.
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                                                                                                                                                                                                                                                                                                                                                                                    PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               STRAIN=ATCC 9790;
MEDLINE=92165737; PubMed=1347040;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       J. Bacteriol. 174:1619-1625(1992)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             STRAIN=ATCC 9790;
MEDLINE=89327152; PubMed=2753858;
                                                                                                                                                                                                                                                                                                                                                                                                                                                           01-FEB-1995 (Rel. 31, Created)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             EMBL; M77639; AAA24776.1; -. PIR; A42296; A42296.
HSSP; P22629; 1SWD.
                                                                                                                                                         :| | |||:| |
462 DL-VNKQDEDPK 472
                                                                                                             200 ELKVLKODKDTK 211
                                                                                                                                                                                                                                                                                                                                                                                    STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Enterococcus hirae."
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     NCBI_TaxID=1354;
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FUNCTION.
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242 TSGNSGGSATT-----TGTTYTVKSGDSV-WGISHSFGITWAQLIEW-NNIKNNFIYPG 293
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTH------VRWFLNINNEKSYVS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   71 KDITIKDQIQGGQQLDLSTLNI------NVTG-THSNYYSGQSAITDFEKAFPGSKITV
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 123 DNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNH
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MEDLINE=97251357; PubMed=9097039;
Aiba H., Baba T., Fujita K., Hayashi K., Inada T., Isono K., Itoh T.,
Asaai H., Kashinoto K., Kimura S., Kitakawa M., Kitagawa M.,
Kasai H., Kashinoto K., Kimura S., Mori H., Mori T., Motomura K.,
Makino K., Miki T., Mizobuchi K., Mori H., Mori T., Motomura K.,
Nakade S., Nakamura Y., Nashimoto H., Nishio Y., Oshima T., Saito N.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   46; Gaps
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STRAINS-RIZ / MGJ652,
STRAINS-RIZ / MGJ652,
STRAINS-9742607; PubMed=9278503;
Blattner F.R., Plunkett G. III, Bloch C.A., Perna N.T., Burland V.,
Riley M., Collado-Vides J., Glasner J.D., Rode C.K., Mayhew G.F.,
Gregor J., Davis N.W., Kirkpatrick H.A., Goeden M.A., Rose D.J.,
Mau B., Shao Y.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Bacteria, Proteobacteria, Gammaproteobacteria, Enterobacteriales,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 666;
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                                                                                                                                                                                                                                                                                                                            Hydrolase; Glycosidase; Bacteriolytic enzyme; Cell wall;
Cell division; Septation; Repeat; Signal.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 1;
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24.0%; Pred. No. 9.4;
tive 28; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                    MURAMIDASE-2
InterPro; IPR002901; Amidase_4.
InterPro; IPR000423; Flag_FlgJ.
InterPro; IPR000432; LysM.
InterPro; IPR000437; Prok_lipoprot_S.
Pfam; PF01832; Amidase_4; 1.
Pfam; PF01875; LysM; 6.
PRINTS; PR01002; FLGFLGJ.
SWART; SM00257; LysM; 6.
SWART; SM00047; LYZZ; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       394 STGNTSSNTANTGSTTSGSTYTVK 417
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LYSM 2.
LYSM 3.
LYSM 4.
LYSM 5.
LYSM 6.
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665
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Bacteria; Firmicutes; Clostridia; Clostridiales; Clostridiaceae;
Clostridium.
  Clostridium botulinum.
                                                                SEQUENCE FROM N.A.
                                      NCBI_TaxID=1491;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ACT SITE
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 This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@libe.sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               82 GQQL----DLSTLNIN-----VTGTHSNYYSGQSAITDFEK------AFPGSK 119
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            214 NGELVITGDNATVNNNGKTTVDGKDSTGTEINGNNGK-VIQDGDLDVSGGGHGIDITGDS 272
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    120 ITVDNTKNTIDVTIPQGYG---SYNSFSINY--KTKITNEQOKEFVNNSQAMYQEHGKEE 174
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               273 ATVDN-KGTMTVTDPESMGIQIDGDKAIVNNEGESTITNGGTGTQINGDDATANNNGKTT 331
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ---STNVTVHKSEA 35
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  36 GTS---SVFYY---KTGDMLPEDTTHVRWFLNI-----NNEKSY--VSKDITIKDQIQG 81
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        84; Gaps
                                                                                                  MEDLINE=92190338; PubMed=1665988;
MOSZET I., Glaser P., Danchin A.;
Moltiple IS insertion sequences near the replication terminus in Escherichia coli K-12.
Biochimie 73:1361-1374(1991).
-! SIMILARITY: TO S.TYPHIMURIUM ORF NEAR CYSG (AC P25928).
-! CAUTION: THIS IS A CONCEPTUAL TRANSLATION; THE GENE CODING FOR THIS PROTEIN IS INTERRUPTED BY A HYBRID IS2D/IS3OC ELEMENT BETWEEN AMINO ACIDS 839 AND 840.
Sampei G., Seki Y., Sivasundaram S., Tagami H., Takeda J., Takemcto K., Takeuchi Y., Wada C., Yamamoto Y., Horiuchi T., "A 570-kb DNA sequence of the Bscherichia coli K-12 genome corresponding to the 28.0-40.1 min region on the linkage map.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 1; Length 2003;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      92; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    2003 AA; 205949 MW; B83A12C8B53220EE CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                            Hypothetical protein; Complete proteome.

CONFLICT 489 489 I -> V (IN REF. 2).

CONFLICT 495 495 I -> V (IN REF. 2).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Pred. No. 37;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              8.5%; Score 95; 25.1%; Pred. No.
                                                                                                                                                                                                                                                                                                                                  EMBL; AE000237; AAC74483.1; ALT SEC.
EMBL; D90778; BAA15009.1; ALT SEQ.
EMBL; D90778; BAA15009.1; ALT SEQ.
EMBL; D90779; BAA18880.1; ALT SEQ.
EMBL; D90779; BAA18880.1; ALT SEQ.
EMBL; X62680; -; NOT ANNOTATED CDS.
ECGENE; EG11307; ydbA.
                                                                [3]
SEQUENCE OF 464-2003 FROM N.A.
                                                   DNA Res. 3:363-377(1996).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       69; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity
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                                                                                                                                                                                                                                                                                                      ENDOPRETIONS:

CATALYTIC ACTIVITY: Limited hydrolysis of proteins of the neuroexocytosis apparatus, synaptobrevins, SNAP25 or syntaxin. No detected action on small molecule substrates:

CHORACTOR: Binds 1 zinc ion per subunit (By similarity).

CHORACTOR: Binds 1 zinc ion per subunit (By similarity).

CHORACTOR: Binds 1 zinc ion per subunit (By similarity).

CHORACTOR: Binds 1 zinc ion per subunit (By similarity).

CHORACTOR: Binds 1 zinc ion per subunit (By similarity).

CHORACTOR: Binds 1 zinc ion per subunit (By similarity).

CHORACTOR: Socreted (By similarity).

CHORACTOR: There are seven antigenically distinct forms of botulinum neurotoxin: Types A, B, Cl. D, E, P, and G.

CHORACTOR: SIMILARITY: Belongs to peptidase family M27.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           96 GTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQGYGS-YNSFSINY-----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   BOTULINUM NEUROTOXIN G, LIGHT-CHAIN.
BOTULINUM NEUROTOXIN G, HEAVY-CHAIN.
ZINC (CATALYTIC) (BY SIMILARITY).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    : 69
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 1296;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    66; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                149013 MW; DC8E47E15F665C31 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ZINC (CATALYTIC) (BY INTERCHAIN (PROBABLE)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 1,
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PROSITE; PS00142; ZINC_PROTEASE; 1.
Neurotoxin; Hydrolase; Metalloprotease; Zinc.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                8.3%; Score 93.5; DE
23.0%; Pred. No. 28;
tive 32; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                BY SIMILARITY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    BY SIMILARITY ZINC (CATALYT)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          InterPro; IPR008985; ConA like lec_gl.
InterPro; IPR002160; Kunitz_legume.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        InterPro; IPR006025; Pept M_Zn_BS.
InterPro; IPR0001395; Peptidase M27.
Pfan; PF01742; Peptidase M27. I
PRINTS; PR00760; BONTOXILXSIN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            EMBL; X74162; CAA52275.1; -. HSSP; P10845; 3BTA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    50, Conservative
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01-NOV-1997 (Rel. 35, Created)
01-NOV-1997 (Rel. 35, Last sequence update)
28-FBE-2003 (Rel. 41, Last annotation update)
Botulinum neurotoxin type G precursor (EC 3.4.24.69) (BONT/G)

BOTG.

1296 AA

STANDARD;

CLOBO

BXG_CLOE Q60393;

GEEDTAN

DNA CLEAVAGE (BY SIMILARITY)

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01-OCT-1993
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 15-DEC-1998
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SEQUENCE
ACT SITE
SEQUENCE
                                                           Query Match
                                                                                  Local
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                                        944 NNNDIQTYLQNEYTIISCIKNDSGWKVSIKGNRIIWTLIDVNAKSKSIFFEYSIKDNISD 1003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         "Complete sequence analysis of the genome of the bacterium Mycoplasma
                                                                                                                                                                                                                                                          01-NOV-1997 (Rel. 35, Created)
01-NOV-1997 (Rel. 35, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
DNA topoisomerase I (EC 5.99.1.2) (Omega-protein) (Relaxing enzyme)
(Untwisting enzyme) (Swivelase).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      SMART; SM00493; TOPRIM; I.
TGRPAMs; TIGROLUSI, topA bact; I.
PROSITE; PS00396; TOPOISOMERASE I PROK; 1.
Isomerase; Topoisomerase; DNA-bInding; Zinc-finger; Metal-binding;
                                                                                                                                                                                                                                                                                                                                                                                                            Bacteria; Firmicutes; Mollicutes; Mycoplasmataceae; Mycoplasma
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Himmelreich R., Hilbert H., Plagens H., Pirkl E., Li B.-C.,
148 -KTKITNEOOKEF----VNNSOAW-YOEHGKE----EVNGKS---
                                                                                                       1004 YINKWFSITITNDRLGNANIYINGSLKKSEKILNLDR 1040
                                                                                ----FNHTVHNI----NANAGIEGTVKGELKVLKQDK 208
                                                                                                                                                                                                                            711 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      InterPro; IPR0005733; DNA_topl_bact.
InterPro; IPR0005733; DNA_tpisomrase.
InterPro; IPR003601; DNA_tpisomrase.
InterPro; IPR003601; DNA_topl_ATP_bind.
InterPro; IPR061571; Toprim_dom.
InterPro; IPR06154; Toprim_sub.
Pfam; PF01131; Topoisom bac; 1.
Pfam; PF01131; Topoisom bac; 1.
Pfam; PF01751; Toprim; 1.
Pfam; PF01396; zf-C4_Topoisom; 2.
PRINTS; PR00417; PRTPISMRASEI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           C4-TYPE 1.
C4-TYPE 2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                    [1]
SEQUENCE FROM N.A.
STRAIN=ATCC 29342 / M129;
MEDLINE=97105885; PubMed=8948633;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AE000056; AAB96220.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SMART; SM00437; TOPIAC; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Complete proteome
                                                                                                                                                                                                                            STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                         Mycoplasma pneumoniae.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PIR; S73898; S73898.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           P06612; 1ECL
                                                                                                                                                                                                                                                                                                                                                                                                                                 NCBI_TaxID=2104;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Herrmann R.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  pneumoniae."
                                                                                                                                                                                                                            TOP1 MYCPN
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Repeat; (ZN_FING ZN_FING
                                                                                                                                                                                                                                             P78032;
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                                                                                                                                                                                                                                                                 494
                                                                                                                                                                                                                                                                                                                                       93 NVTGTH-SNYYSGQSAITDFEKAFPGSKITVDNTKNTI-DVTIPQGYGSYN--SFSINYK 148
                                                                                                                                                                                                                                                                                                                                                                                                      495 TARQTHPAARYTQASLIEALEK----SNIGRPSTYNTMASVNLDRGYASLNKHAFHVTQL 550
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        149 TKITNEQ------QKEFVNNSQAWYQEHGKEEV--NGKSFNHTVHNINANAGIE-GT 196
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  436 THIR-FINDGNKFYASSKSLVFDGYRKIYEHFENKESNDLYIDLDKIRVGDRFMAKDIKI
                                                                                                                                                                                               54 THVRWFLNINNE-----KSYV-----SKDITIK-DQIQGGQLDLSTLNI
                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Bhat K.S., Gibbs C.P., Barrera O., Morrison S.G., Jaehnig F., Stern S., Kupsch B.-M., Meyer T.F., Swanson J.; Mol. microbiol. 6:1073-1076(1992).
-!- FUNCTION: IMPLICATED IN A NUMBER OF ADHERENCE FUNCTIONS. OPA PROTEINS ARE IMPLICATED IN PATHOGENESIS AND ARE SUBJECT TO PHASE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MEDLINE=93178439; PubMed=8440254;
Kupsch B.-M., Knepper B., Kuroki T., Heuer I., Meyer T.F.;
"Variable opacity (Opa) outer membrane proteins account for the cell
"Topisms displayed by Neisseria gonorrhoeae for human leukocytes and
                                                                                                                                    47;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MEDLINE=9211/767; PubMed=1815562; Merrison S.G., Jaehnig F., Bhat K.S., Gibbs C.P., Barrera O., Morrison S.G., Jaehnig F., Stern A., Kupsch E.-M., Meyer T.F., Swanson J.; "The opacity proteins of Neisseria gonorrhoeae strain MS11 are encoded by a family of 11 complete genes."; Mol. Microbiol. 5:1889-1901(1991).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Bacteria, Proteobacteria, Betaproteobacteria, Neisseriales,
                                                                                                                                Indels
                                                                     Length
711 AA; 81965 MW; 16B863588B06582E CRC64;
                                                                                                                                65;
                                                                  DB 1,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  238 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Opacity protein OPA60 precursor (Fragment).
                                                              8.2%; Score 92.5; D
25.4%; Pred. No. 16;
iive 32; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                SUBCELLULAR LOCATION: Outer membrane.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MEDLINE=92261323; PubMed=1584024;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Created)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            197 VKGELKVLKQDKD 209
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   :|::|:
606 AEGSIQRVKKEKE 618
                                                                                                                                    49; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Neisseriaceae; Neisseria.
NCBI_TaxID=485;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        EMBO J. 12:641-650(1993).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (Rel. 27, (Rel. 27,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Neisseria gonorrhoeae
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (Rel. 37,
                                                                                                 Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  204 HKQWGSSGKINYLKEAQNMITKGIKASNVTKNNGLNLGDWGDKSTF-----DTRPSDWMM 258
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         298 İSDEVVKNPPQPAPKDFLDESKYTDSYYYNASRVPLRİVMDYAMYGEKRGKVISDKVATW 357
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    145 INYKTK------ITNEQOKEFVNNSQAWYQEHG 171
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         358 IKSKTKGNPSKIVDGYKLDGTNIGDYPTAVYVSPFIAAGTTNSKNQEWVNSGWDW-MKNK 416
                                                                                                                                                                                                                                                                                                                                                                                                                                                     7 HHHHGSDDKV-----ATITSGNKSTNVT----VHKSEAGTSSVFYYKTGDMLPED--T
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      54 THVRWFLNINNEKSYVSKDITIKDQIQGGQLDLSTLNI--NVTGTHSNY---YSGQSA-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        STRAIN=UAB CTIP;
MEDLINE=21267165; PubMed=11353084;
Chambaud I., Heilig R., Ferris S., Barbe V., Samson D., Galisson F.,
Moszer I., Dybvig K., Wroblewski H., Viari A., Rocha E.P.C.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Shen X., Gumulak J., Yu H., French C.T., Zou N., Dybvig K.; "Gene rearrangements in the vsa locus of Mycoplasma pulmonis."; J. Bacteriol. 182:2900-2908(2000).
                                                                                                                                                                                                                                                                                                                                                                                                                      64; Indels 102;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Bacteria; Firmicutes; Mollicutes; Mycoplasmataceae; Mycoplasma.
                                                                                                                                                                                                                                                                                                                                                                                Score 91, DB 1; Length 463;
Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      108 IIDF---EKAFPGSKITVDNTKNT-----IDVTIPQGYGSYN---
                                                                                                                                                                                                                                                                                 POTENTIAL.
ENDOGLUCANAGE.
PROTON DONOR (BY SIMILARITY)
WUCLEOPHILE (POTENTIAL).
407FAS4F5236C59E CRC64;
                                                                                                                                                                                                                                 PROSITE; PS00812; GLYCOSYL_HYDROL_F8; 1.
Cellulose degradation; Hydrolase; Glycosidase; Signal.
SIGNAL
PROPEP
CHAIN
S6 463 ENDOGLORANSE.
CHAIN
S6 463 ENDOGLORANSE.
ACT SITE
130 130 PUCLEOPHILE (POTENTIAL).
ACT SITE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       050274; 098039;
16-0CT-2001 (Rel. 40, Created)
16-0CT-2001 (Rel. 40, Last sequence update)
10-0CT-2003 (Rel. 42, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         578 AA
                                                                                                                                                                                                                                                                                                                                                                                                                    31; Mismatches
                                                                                                                               EMBL; M68872; AAA22409.1; -.
PIR; A44808; A44808.
InterPro; IPR002037; Glyco.hydro.8.
InterPro; IPR008928; Glyco_trans_6hp.
Pfam; PF01270; Glyco.hydro.8; 1.
PRINTS; PR00735; GLH\(T\)DRLASE8.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          STRAIN=KD735-15;
MEDLINE=20245550; PubMed=10781561;
                                                                                                                                                                                                                                                                                                                                                     51882 MW;
                                                                                                                                                                                                                                                                                                                                                                                     8.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                      53; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Lipoprotein A precursor.
LIPA OR MYPU_5300.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       172 KEEVNGKSFN 181
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      417 KESYFSDSYN 426
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Mycoplasma pulmonis.
                                                                                                                                                                                                                                                                                                                                  191 1
463 AA;
                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
Matches 53; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                      -KAFPGSKI-----TVDNTKNTIDV-TIPQG--YGSYNSFSINYKTKITNEQQKEFVN 161
                                                                                                                                                                                                                                                                                                                                                 16 AYAYEHITHDYPEPTAPNKNKISTVSDYFRNIRTRSVHPRVSVGYDFGGWRIAADYARYR 75
                                                                                                                                                                                                                                                                                                                                                                                     -WFLNINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFE---
                                                                                                                                                                                                                                                                                                                      SHHHHHHGSDDKVATITSGNKSTNVTVHKSEAGTSSVF----YYKTGDMLPEDTTHVR
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       PH AT 5.2.
-!-CATALYTIC ACTIVITY: Endohydrolysis of 1,4-beta-D-glucosidic
linkages in cellulose, lichenin and cereal beta-D-glucans.
-!- PTM: THE N- AND THE C-TERMINUS MAY BE SUBJECTED TO PROTEOLYSIS.
-!- MISCELLANEOUS: ONE TRP RESIDUE HAS BEEN PROVED TO BE INVOLVED IN
-!- SIMILARITY: Belongs to cellulase family D (family 8 of glycosyl hydrolases).
                                                                                                                                                                                                                                                                                       Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Ozaki K., Ito S.;
"Purification and properties of an acid endo-1,4-beta-glucanase from
Bacillus sp. KSM-330.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Gen. Microbiol. 137:41-48(1991).
- FUNCTION: THIS ACID ENDOGLUCANASE IS ACTIVE OVER AN EXTREMELY
NARROW RANGE OF PH VALUES, BETWEEN 4.5 AND 6.5, WITH AN OPTIMUM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          gene encoding an
                                                                                                                                                                                                                                                                                     33;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    01-DEC-1992 (Rel. 24, Created)
01-DEC-1992 (Rel. 24, Last sequence update)
28-FEB-2003 (Rel. 41, Last annotation update)
Endoglucanase precursor (EC 3.2.1.4) (Endo-1,4-beta-glucanase)
                                                                                                                                                                                                                                                      DB 1; Length 238;
                                                                                                                                                                     / V18).
                                                                                                                                                                                                                                                                                     Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (Cellulase) (Endo-K).
Bacillus sp. (strain KSM-330).
Bacteria, Firmicutes; Bacillales; Bacillaceae; Bacillus.
                                                                                                                                                                                                                     27073 MW; 883A3560C2DF1B9F CRC64;
                                                                                                                                  POTENTIAL.
OPACITY PROTEIN OPA60.
SED -> MLKA (IN MS11 / V
V -> M (IN MS11 / V18).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Ozaki K., Sumitomo N., Ito S.; Molecular cloning and nucleotide sequence of the endo-1, 4-beta-glucanase from Bacillus sp. KSM-330. J. Gen. Microbiol. 137:2299-2305(1991).
                                                                                                                                                                                                                                                                                     69
                                                                                                                                                                                                                                                                  4.7;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        463 AA.
               EMBL; Z18939; CAA79372.1; -.
EMBL; X60711; CAA43121.1; -.
INFR; S16619; S16619; Dorin_opacity.
Pfam; PF02462; Opacity; 1.
Outer membrane; Multigene family; Signal.
SIGNAL <1 1 POTENTIAL.
                                                                                                                                                                                                                                                                                     Conservative 35; Mismatches
                                                                                                                                                                                                                                                    8.2%; Score 91.5;
24.7%; Pred. No. 4.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SEQUENCE OF 56-75, AND CHARACTERIZATION MEDLINE=91259037; PubMed=2045781;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        STANDARD;
                                                                                                                                                    >238
                                                                                                                                                                                   234
                                                                                                                                                                                                                   238 AA;
                                                                                                                                                                                                                                                                Local Similarity
es 45; Conserv
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SEQUENCE
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STRAIN=HD-525
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Q8MTI2;
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                                                                                                                                                                                                                                                                                                                                                               This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation-the European Bioinformatics Institute. There are no restrictions on its most by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        121 IQENKQ----SQKDQNTSAVNVSALE-----KQTKNDENISLVNSKDTNVILKNDEKVA 170
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        231 LSLTFKEGMPLGLEVVLKLENLDSHEEKEISFKTTNGKVQNVLLTSSNLTSGKWKIKSFS 290
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Lipoprotein A.

N-palmitoyl cysteine (Potential).
S-diacylglycerol cysteine (Potential).
ASN-RICH.

K -> R (IN STRAIN KD735-15).
N -> D (IN STRAIN KD735-15).
N -> D (IN STRAIN KD735-15).
S -> A (IN STRAIN KD735-15).
Y -> N (IN STRAIN KD735-15).
Y -> N (IN STRAIN KD735-15).
WH; BS33ED3467005D89 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      73 ITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           171 LAKDDSKEKSKNSSNLNLKTPVENRONKNEVKDDKKALOWWOKLNESASILESFSYDQTS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                14 DKVATITSGNK-STNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKD
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                133 IPQGYG---SYNSFSINYKTKITNEQQKEFVNNSQ---AWYQE-------
                                                                                                                                                                           BLUGTE B., VOELKET L.L., ZOU N., YU H., Dybvig K.;
"Mechanism of antigenic variation in Mycoplasma pulmonis: interwoven, site-specific DNA inversions.";
Mol. Microbiol. 18:703-714(1995).
-!- SUBCELLULAR LOCATION: Attached to the membrane by a lipid anchor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                   (Potential).
-!- SIMILARITY: BELONGS TO THE PULMONIS LIPAB LIPOPROTEIN FAMILY.
                        complete genome sequence of the murine respiratory pathogen
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Mypulist, MrPU 5300;
-.
Mypulist, MrPU 5300;
InterPro; IPR000437; Prok lipoprot S.
InterPro; IPR000437; Prok lipoprot S.
Pfam; PF04200; Lipoprotein 17; 1.
PROSITE; PS00013; PROKAR LIPOPROTEIN; 1.
Lipoprotein; Membrane; Signal; Complete proteome; Palmitate.
27 Signal; POTENTIAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 1; Length 578;
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                                             Mycoplasma pulmonis.";
Nucleic Acids Res. 29:2145-2153(2001).
                                                                                                                                       STRAIN=KD735-15;
MEDLINE=96414471; PubMed=8817492;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    EMBL; U23947; AAB41030.2; -.
EMBL; AL445565; CAC13703.1; -.
PIR; B90578; B90578.
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                                                                                                              PARTIAL SEQUENCE FROM N.A.
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Best Local Similarity
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Blanchard A.;
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CIGB BACTZ
ID CIGB BACTZ
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PRT; 1169 AA.

STANDARD;

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30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
16-CCT-2001 (Rel. 40, Last annotation update)
Pesticidial crystal protein crylGb (Insecticidal delta-endotoxin
CrylG(b)) (Crystaline encomocidal protoxin) (133 kDa crystal protein).
CRYLGB OR CRYLG(B) OR CRYHZ.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  EPITHELIAL CELLS OF LEPIDOPTERAN LARVAE. TOXIC TO PIERIS RAPAE.
-!- DEVELOPMENTAL STAGE: The crystal protein is produced during sporulation and is accumulated both as an inclusion and as part of
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     1021 VTAYKEGYGEGCVTIHEI-----EDNTDELKFSNCVEEEIYPNNTVTCNDYTATQEEYE-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             28 VIVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIKDQIQGQQLDL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       wuhanensis strain.";
Curr. Microbiol. 40:227-232(2000).
-!- FUNCTION: PROMOTES COLLOIDOSMOTIC LYSIS BY BINDING TO THE MIDGUT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 13; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      the spore coat.
-!- MISCELLANEOUS: Toxic segment of the protein is located in the N-
                                                                                                                                                                                                                                                                                                                                                                                                                           Kuo W.-S., Lin J.-H., Tzeng C.-C., Kao S.-S., Chak K.-F.; "Cloning of two new cry genes from Bacillus thuringiensis subsp.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 1169;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  1169 AA; 132904 MW; DIEFC1508A8B10BD CRC64;
                                                                                                                                                                                        Bacillus thuringiensis (subsp. wuhanensis).
Bacteria; Firmicutes; Bacillales; Bacillaceae; Bacillus.
NCBI_TaxID=52024;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             SIMILARITY: Belongs to the delta endotoxin family.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           LS-WAR-2004 (Rel. 43, Created)
15-WAR-2004 (Rel. 43, Last sequence update)
15-WAR-2004 (Rel. 43, Last annotation update)
15-WAR-2004 (Rel. 43, Last annotation update)
15-WAR-2004 (Rel. 43, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              8.1%; Score 90.5; DE 24.5%; Pred. No. 41; rative 17; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     625 AA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        148 KTKITNEQQKEFVNNSQAWYQEHGKEE 174
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              InterPro; IPR005638; endotoxin_C.
InterPro; IPR005639; endotoxin_N.
InterPro; IPR008979; Gal_bind_like.
Pfam; PF00355; endotoxin_C; 1.
Pfam; PF03944; endotoxin_C; 1.
Pfam; PF03945; endotoxin_C; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PRT;
                                                                                                                                                                                                                                                                                                                                                                                      MEDLINE=20153386; PubMed=10688690;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       InterPro; IPR001178; Endotoxin.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 EMBL; U70725; AAD10291.1; -.
HSSP; P02965; 1CIY.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Toxin; Sporulation.
SEQUENCE 1169 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Local Similarity
                                                                                                                                                                                                                                                                                                                          SEQUENCE FROM N.A.
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ID BSL1_TT
AC Q8MT12
DT 15-MAR
DT 15-MAR
DE PUTALI
GN BSPAL1
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation. The European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (see http://www.isb-sib.ch/announce) or send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 232 DSVTTIANSAFYECSKLTSITIGKSVTRIEGNAFSKCYSLTSITIKTTNDITSSITTDV- 290
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       291 -FLNCPITELIYETTGLTFLTYEYFKDKV------TLIKFNIPKSDSNSMIRLQEITS 341
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   105 QSAITDFEKAFPGSKITVDNTKNTIDVTIPQGY--------GSYNSFSI 145
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    :| | : |:|:: | ::||342 LPTLTHFTNL---NKVTIENIN---ELTIPESFIEGDNFEILITWNIKSIDPNAFKDCSI 395
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  58 WFLNIN-NEKSYVSKDIT-----IKDQIQGQQLDLSTLNINVTGTHSN-----YYSG 104
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match 8.0%; Score 90; DB 1; Length 625; Best Local Similarity 19.9%; Pred. No. 20; Matches 46; Conservative 42; Mismatches 73; Indels 70; Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   14 DKVATITSG-----NKSTNVTVHKSEA------GTSSVFYYKTGDMLPEDTTHVR
                                                                           MEDLINE=22356757; PubMed=12467987;
MEDLINE=22356757; PubMed=12467987;
Hirt R.P., Harriman N., Kajava A.V., Embley T.M.;
"A novel potential surface protein in Trichomonas vaginalis contains a leucine-rich repeat shared by micro-organisms from all three
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   NYKTKI-TNEQQKEFVNNSQAWYQ-----EHGKEEVNGKSFNHTVHNINAN 190
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     396 NKFTYLGTDKLENDFLKNAKSCEEVITSTKYSDNEIGGMITHKQSEENPN 446
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               N-LINKED (GLCNAC. . .) (POTENTIAL)
N-LINKED (GLCNAC. . .) (POTENTIAL)
62EF37BDA8AD001A CRC64;
            Eukaryota; Paržbasalidea; Trichomonadida; Trichomonadidae;
Trichomonadinae; Trichomonas.
NCBI_TaxID=5722;
                                                                                                                                                                                                                                                                                                                                                                                                                                Leucine-rich repeat.
EXTRACELLULAR (POTENTIAL).
POTENTIAL.
                                                                                                                                                                           MOI. Blochem. Parasitol. 125:195-199 (2002).
-1- FUNCTION: May bind host tissue.
-1- SUBCELLULAR LOCATION: Surface membrane (Probable).
-1- SIMILARITY: Contains 13 leucine-rich (LRR) repeats.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              CYTOPLASMIC (POTENTIAL)
LRR 1.
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LER 5.
LER 6.
LER 7.
LER 9.
LER 10.
LER 11.
LER 11.
LER 11.
LER 12.
LER 13.
ASN/PRO-RICH.
                                                                                                                                                                                                                                                                                                                                                                                              EMBL; AY101349; AAM51159.1; -. InterPro; IPR007093; LRR_Tp.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   67021 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                              Repeat; Transmembrane;
DOMAIN 1 548
Trichomonas vaginalis.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  15
227
625 AA;
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Q9F865 Q9F3612 Q9F362 Q9F360 Q9FB60 Q9FB60 Q9FB7 Q9FBV5 Q9FBV5 Q9FBV5 Q9FBV5 Q8FBV5 Q8FBV5 Q8FBV7 Q8FBV7 Q8FBV7 Q8FBV7 Q8FBV7 Q8FBV7 Q8FBV7 Q8FBV7

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SEQUENCE FROM N.A.
MEDLIKES-22040717; PubMed=12044378;
MEDLIKES-22040717; PubMed=12044378;
MEDLIKES-22040717; PubMed=12044378;
MEDLIKES-22040717; PubMed=12044378;
Nagai Y., Takeuchi F., Kuroda M., Asano K., Naimi T., Kuroda H., Cui L.,
Yamamoto K., Hiramatsu K.;
Menome and virulence determinants of high virulence community-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1183 AA; 133001 MW; 7A33DA876F9DA447 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-OCT-2002 (TrEMBLrel. 22, Created)
01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
Collegen adhesin.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  92.2%; Score 1035; DB 16; 98.5%; Pred. No. 7.6e-58; ive 1; Mismatches 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Staphylococcus aureus (strain MW2).
Bacteria; Firmicutes; Bacillales; Staphylococcus.
NCBI_TaxID=196620;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Lancet 359:1819-1827 (2002).

EMBL; AP004831; BAB96477.1; -
PIR; A44404; A42404.

PIR; A446204; A48620.

InterPro; IPR008956; Adhes_bact.

InterPro; IPR008956; Cala B. unit.

InterPro; IPR008456; Calagen_bind.

InterPro; IPR008456; Collagen_bind.

Pfam; PF05738; Cal B; 7.

Pfam; PF05738; Callagen_bind; 1.
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Complete protecme
SEQUENCE 1183 AA; 133001 MW; 7A331
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Best Local Similarity 98.5
Matches 197; Conservative
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acquired MRSA.";
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Q840a5 streptococc
Q8kr25 streptococc
Q7wwp3 arcanobacte
Q9f865 enterococcu
Q9f866 enterococcu
Q9f867 enterococcu
Q9f877 enterococcu
Q9f877 enterococcu
Q9f862 enterococcu
Q9f864 enterococcu
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1707.034 Million cell updates/sec
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1122
1 MRGSHHHHHHGSDDKVATIT......GIBGTVKGBLKVLKQDKDTK 211
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                                                                                                            July 27, 2004, 10:05:01 ; Search time 39 Seconds
                                                                                                                                                                                                                                                                                                                       1017041
                GenCore version 5.1.6
Copyright (c) 1993 - 2004 Compugen Ltd.
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                                                                                                                                                                                                                                                                                        1017041 seqs, 315518202 residues
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Maximum Match 100%
Listing first 45 summaries
                                                                                - protein search, using sw model
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sp_rodent:*
sp_virus:*
sp_vertebrate:*
sp_unclassified:*
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Q9F861
Q9F864
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Gapop 10.0 , Gapext 0.5
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sp bacteria: *
sp fungi: *
sp human: *
sp nammal: *
sp mammal: *
sp organele: *
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sp_bacteriap:*
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Maximum DB seq length: 200000000
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Match Length DB
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No.
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PRT; 1183 AA

ALIGNMENTS

Q8XP11 Q86ZE8 Q86ZE8 Q816Z Q931E7 Q931E7 Q95EPH4 Q813E5 Q813E7 Q813E7 Q81C27

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 18 TITSGNKSTNVTVHKSEAGT-SSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIK
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Submitted (JUN-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AF521599; AAM77657.1; -.
InterPro; IPR008966; Adhes bact.
InterPro; IPR008456; Collagen bind.
Pfam; PF05737; Collagen bind.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   41.3%; Score 463.5; DB 2; Length 657;
47.7%; Pred. No. 1e-21;
tive 36; Mismatches 62; Indels 5;
                                                                                                                                                                                                                                                                             Lannergard J., Frykberg L., Guss B.,
Lannergard J., Frykberg L., Guss B.,
"CNE, a collagen-binding protein of Streptococcus equi.";
Submitted (DEC-2002) to the EMBL/GenBank/DDBJ databases.
EMBL, AX193773, AA043099.1; -.
InterPro; IPR008966; Adhes_bact.
InterPro; IPR008966; Adhes_bact.
InterPro; IPR008999; Gram_pos_anchor.
IIGRFAMS; TIGR01167; LPXTG_anchor; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-0CT-2002 (TrEMBLrel. 22, Created)
01-0CT-2002 (TrEMBLrel. 22, Last sequence update)
01-0CT-2003 (TrEMBLrel. 25, Last annotation update)
Putative collagen adhesin se73.9 (Fragment).
Streptococcus equi.
Bacteria, Firmicutes; Lactobacillales; Streptococcaceae;
                                                                                                            Bacteria; Firmicutes; Lactobacillales; Streptococcaceae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    657 657
657 AA; 73889 MW; A902E85BBCE59C4F CRC64;
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221 AA; 25049 MW; 0300F430D21EB859 CRC64;
01-OCT-2003 (TrEMBLrel. 25, Last annotation update) Collagen-binding protein Cne precursor (Fragment).
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                                                                                  Streptococcus equi subsp. equi
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              197 -- VKGELKVLKQDKDTK 211
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             94; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Local Similarity
                                                                                                                                                                    NCBI_TaxID=148942;
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                                                                                                                                                                                                                                                            STRAIN=1866;
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QBKRZ5;
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                             DITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDV 131
                                                                                                                                                                                                                                                            264 TIPQGYGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANA 323
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                                                                                                                                            132 IIPQCYGSYNSFSINYXTXITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANA
  SDDKVATITSGNKSTNVTVHKSEAGTSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSK
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Enterococcus faecium (Streptococcus faecium).
Bacteria, Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STRAIN=TX2555;
MEDLINE=22511562; PubMed=12622825;
Mallapareddy S.R., Weinstock G.M., Murray B.E.;
"Clinical isolates of Enterococcus faecium exhibit strain-specific collagen binding mediated by Acm, a new member of the MSCRAWM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    721 AA; 81545 MW; DB5BB3B3DEF45673 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        01-JUN 2003 (TrEMBLrel. 24, Created)
01-JUN-2003 (TrEMBLrel. 24, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
Collagen adhesin
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Q840AS PRELIMINARY; PRT; 657 AA. Q840AS; (TrEMBirel. 24, Created) 01-JUN-2003 (TrEMBirel. 24, Last sequence update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         721 AA
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EMBL, AY135217; AAN12397.1; -.

InterPro; IPR008966; Adhes bact.

InterPro; IPR008990; Gram B_unit.

InterPro; IRR01899; Gram_Dos anchor.

Pfam; PP00746; Gram pos anchor; 1.

PROSITE; PS50847; GRAM_POS_ANCHORING; 1.
                                                                                                                                                                                                                                                                                                                                           192 GIEGTVKGELKVLKQDKDTK 211
                                                                                                                                                                                                                                                                                                                                                                        196 TVKGELKVLKQDKDTK 211
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|TVRGELKIFKYINDTE 345
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Best Local Similarity
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SEQUENCE
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SEQUENCE FROM N.A.

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SEQUENCE FROM N.A.

SEQUENCE STRAIN-Beirut/TX0645;

MEDIINE=20407335; PubMed=10948146;

Mallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;

Tolerand and ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enterococcus faecalis and evidence for production of ace during human infect: Immun. 68:5210-5217(2000).

Infect: Immun. 68:5210-5217(2000).

REMBL; AF260889; AAG23847.1; -..

RHSSP; Q53654; IAMX.

InterPro; IPR008456; Adhes bact.

RINTERPRO; IPR008456; Collagen_bind.

RINTERPRO; IPR008456; Collagen_bind.

PUNN TER 319 319
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    171 JADRQGSGQQLINKESFTFDIVNDKETKY---ISLAEFEQOGYGKIDFV--TDNDFNLRFY 225
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                226 QDKARFTSFIVRYTSTITEACGHQATFENSYDINYQLMNQDATNEKNTSQV-----KNVF 280
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    135 QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
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STRAINE-END6/TX0045;
MEDLINE=20467335; PubMed=10948146;
Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
"Diversity of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enterococcus faecalis and evidence for production of ace during human
                                                        Enterococcus faecalis (Streptococcus faecalis).
Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
NCBI_TaxID=1351;
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Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
NCBI_TaxID=1351;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ; Score 182; DB 2; Length 319;
; Pred. No. 0.00039;
32; Mismatches 88; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        319 AA; 35986 MW; DSF914E0856A6940 CRC64;
       01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
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Last annotation update)
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InterPro; IPR008456; Collagen bind.
InterPro; IPR001899; Gram pos_anchor.
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EMBL; AF260873; AAG23932.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity 28.8%;
Matches 55; Conservative 3:
                            Collagen adhesin (Fragment).
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01-MAR-2001 (TrEMBLrel. 16,
01-OCT-2003 (TrEMBLrel. 25,
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MEDLINE=2275633; PubMed=12874314;

MEDLINE=2275633; PubMed=12874314;

Esmay P.A., Billington S.J., Link M.A., Songer J.G., Jost B.H.;

Esmay P.A., Billington S.J., Link M.A., Songer J.G., Jost B.H.;

The Accanobacterium pyogenes Collagen-Binding Protein, CbpA, Promotes Adhesion to Host Cells.";

Adhesion to Host Cells.";

Infect. Immun. 71:4368-4374(2003).

EMBL, AY223543; AA043108.1;

Collagen; Signal.
                                                                                                   TITSGNKSTNVTVHKSEAGT-SSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDITIK
                                                                                                                             DQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIPQG
                                                                                                                                                                                                                                                                                                137 YGSYNSFSINYKTKITNEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAGIEGT
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     Length 221;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Arcanobacterium pyogenes.
Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Actinomycineae; Actinomycetaceae; Arcanobacterium.
NCBI_TaxID=1661;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Match 16.3%; Score 182.5; DB 2; Length Local Similarity 29.5%; Pred. No. 0.0016; les 61; Conservative 34; Mismatches 91; Indels
                                                   Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   27 Potential.
124719 MW; 3127352B9795CE98 CRC64;
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Last annotation update)
41.2%; Score 462.5; DB 2;
llarity 47.7%; Pred. No. 3.4e-22;
Conservative 36; Mismatches 62;
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01-MAR-2001 (TrEMBLrel. 16,
01-MAR-2001 (TrEMBLrel. 16,
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                            Similarity
                                                   94;
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  Query Match
Best Local 8
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QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
                                                                            266 RNKARPTSFIVRYTSTITEAGQHQATFENSYDINYQLANQDATNEKNTSQV-----KNVF 320
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Enterococcus faecalis (Streptococcus faecalis).
Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
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Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B "Diversity of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enterococcus faecalis and evidence for production of ace during hifections.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              16.0%; Score 179; DB 2; Length 721; 28.3%; Pred. No. 0.0015; ive 33; Mismatches 88; Indels
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721 AA; 78842 MW; 1CA8A1D98CE69392 CRC64;
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1-MAR-2001 (TrEMBLrel. 16, Last sequence update)
01-0CT-2003 (TrEMBLrel. 25, Last annotation update)
Collagen adhesin precursor (Fragment).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (TrEMBLrel. 16, Created)
(TrEMBLrel. 16, Last sequence update)
(TrEMBLrel. 25, Last annotation update)
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InterPro; IPR008456; Collagen_bind.
InterPro; IPR01899; Gram_pos_anchor.
Pfam; PF05737; Collagen_bind; 1.
Pfam; PF07746; Gram_pos_anchor; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Infect. Immun. 68:5210-5217(2000).
EMBL; AF260872; AAG23931.1; -.
HSSP; Q53654; 1AMX.
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321 VEGEASGNONV 331
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              135 QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
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STRAIN=MD9/TX0249;
MEDLINE=20407335; PubMed=10948146;
Mallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
"Diversity of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enterococcus faecalis and evidence for production of ace during human
                                                                                                                                                                                                                                                                                                       Gaps
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Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
NCBI_TaxID=1351;
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                                                                                                                                                                                                                                        Length 721;
                                                                                                                                                                                                                                  Ouery Match 16.1%; Score 181; DB 2; Length 72 Best Local Similarity 28.8%; Pred. No. 0.0012; Matches 55; Conservative 32; Mismatches 88; Indels
                                                                                                                                       31 POTENTIAL.
79047 MW; F09EDBE49E5DF866 CRC64;
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Last annotation update)
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EMBL; AF260877; AAG23935.1; -.
HSSP; Q5364; 1AMX.
INCERPO; IPR008966; Adhes bact.
InterPro; IPR008456; Collagen_bind.
InterPro; IPR001899; Gram pos anchor.
Pfam; PP05737; Collagen_bind; I.
Pfam; PP07746; Gram pos anchor.
Pfam; PF00746; Gram pos anchor; I.
PROSITE; PS50847; GRAM_POS_ANCHORING; 1.
Pfam; PF05737; Collagen bind; 1.
Pfam; PF00746; Gram pos anchor; 1.
PROSITE; PS50847; GRAM POS ANCHORING; 1.
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01-MAR-2001 (TrEMBLrel. 16,
01-OCT-2003 (TrEMBLrel. 25,
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                                                                                                                                                                   721 AA;
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STRAIN=B-143/TXX31;
MEDLINE=20407335; PubMed=10948146;
MEDLINE=20407335; PubMed=10948146;
Mallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
Mallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
The office of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of recognizing adhesive matrix molecules, from different strains of infections.";
The enterococcus faecalis and evidence for production of ace during human infect. Immun. 68:5210-5217(2000).

REMBL; AF260896; AAG23954.1; -.

In fect. Immun. 68:5210-5217(2000).

REMBL; AF260896; AAdhes_bact.

InterPro; IPR008966; Adhes_bact.

RICEPPRO; IPR008966; Adhes_bact.
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Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
NCBI_TaxID=1351;
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                                                                                                                          319 319 36001 MW; 2054C2B11ECC5920 CRC64;
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01-MAR-2001 (TrEMBLrel. 16, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
EMBL; AP260894; AAG23952.1; -. HSSP; 03564; 1AMX. InterPro; IPR008966; Adhes bact. InterPro; IPR008456; Collagen_bind. Ffam; PP05737; Collagen_bind; 1. NON_TER 319 319
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Matches 54; Conserv
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                                                                                                   SEQUENCE FROM N.A.
STRAINE-Various strains;
MEDLINE=20407335; PubMed=10948146;
Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
"Diversity of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enteroccus faecalis and evidence for production of ace during human
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MEDLINER-20407335; PubMed=10948146;
Mallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
"Diversity of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enterococcus faecalis and evidence for production of ace during human infections.";
                       Enterococcus faecalis (Streptococcus faecalis).
Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
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Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
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01-WAR-2001 (TrEMBLrel. 16, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
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EMBL; AF260895; AAG23953.1; --
EMBL; AF260880; AAG23938.1; --
EMBL; AF260881; AAG23940.1; --
EMBL; AF260882; AAG23940.1; --
EMBL; AF260888; AAG23946.1; --
EMBL; AF260898; AAG23946.1; --
EMBL; AF260898; AAG23948.1; --
EMBL; AF260893; AAG23948.1; --
EMBL; AF260893; AAG23948.1; --
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InterPro, IPR008456; Collagen bind
Pfam; PF05737; Collagen_bind, 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Q53654; 1AMX
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                                                           NCBI_TaxID=1351;
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Local Similarity
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        226 RDKARFTSFIVRYTSTITEAGGHQATFENSYDINYQLNNQDATNEKNTSQV----KNVF 280
                                                                                                                                                                                                                                                                                                                                      75 IKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIP 134
                                                                                                                                                                                                                                                                                                                                             135 QGYGSYNSFSINYKTKIT--NEQOKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
                                                                                                                                                                                                                                                                                                                                                                         18 TITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDIT 74
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Enterococcus faecalis (Streptococcus faecalis).
Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
                                                                                                                                                                Length 458;
                                                                                                                                                                                                                                                                                           Indels
                                                                                                                                                                                                                                                              51085 MW; B674692BD29A0A24 CRC64;
                                                                                           01-NOV-1999 (TrEMBLrel. 12, Created)
01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
01-CCT-2003 (TrEMBLrel. 25, Last annotation update)
Collagen adhesin Ace (Fragment).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Last sequence update)
Last annotation update)
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Best Local Similarity 28.3%; Pred. No. 0.0011;
Matches 54; Conservative 33; Mismatches 88
                                                                              458 AA
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                                                                              PRELIMINARY;
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                            193 IEGTVKGELKV 203
                                    458 AA;
                                                                                                                                                          SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             135 QGYGSYNSFSINYKTKIT--NEQQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               266 RDKARFTSFIVRYTSTITEAGQHQATFENSYDINYQLNNQDATNEKNTSQV-----KNVF 320
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              74
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       18 TITSGNKSTNVTVHKSEAG---TSSVFYYKTGDMLPEDTTHVRWFLNINNEKSYVSKDIT
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Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
STRAIN=LBJ-1/TX0020;
MEDLINE=20(907335; PubMed=10948146;
MEDLINE=20(907335; PubMed=10948146;
Mallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.
"Diversity of ace, a gene encoding a microbial surface component recognizing adhesive matrix molecules, from different strains of enterococcus faecalis and evidence for production of ace during human infections.";
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ch 15.9%; Score 178; DB 2; Length 580; Similarity 28.3%; Pred. No. 0.0014; 54; Conservative 33; Mismatches 88; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                POTENTIAL.
7CD61EAD3FDA0993 CRC64;
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Last annotation update)
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Signal. 1 31 POTENTIAL.
                                                                                                                                                                                                                                                                                                                                                                                                                            Pfam; PF05737; Collagen bind; 1.
Pfam; PF00746; Gram pos_anchor; 1.
PROSITE; PS50847; GRAM_POS_ANCHORING; 1.
                                                                                                                                                                                                                             Infect. Immun. 68:5210-5217(2000).
EMBL; AFF560878; AAG23936.1; -.
HSSP; Q53654; LAMX.
InterPro; IPR008966; Adhes bact.
InterPro; IPR008456; Collagen bind.
InterPro; IPR001899; Gram pos_anchor.
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InterPro; IPR008966; Adhes_bact.
InterPro; IPR008956; Collagen bind.
InterPro; IPR001899; Gram_pos_anchor.
Pfam; PP05737; Collagen bind; I.
Pfam; PF00746; Gram_pos_anchor; 1.
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EMBL; AF260879; AAG23937.1; -.
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580 AA; 63948 MW;
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75 IKDQIQGGQQLDLSTLNINVTGTHSNYYSGQSAITDFEKAFPGSKITVDNTKNTIDVTIP 134
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                        Query Match 15.9%; Score 178; DB 2; Length 580;
Best Local Similarity 28.3%; Pred. No. 0.0014;
Matches 54; Conservative 33; Mismatches 88; Indels 16; Gaps
580 AA; 64176 MW; 060EFE50D61A5271 CRC64;
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